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(71) Applicant (for all designated States except AU BB CA G MN MW NZ SD US): UNILEVER N.V. [NL/NL] 455, NL-3013 AL Rotterdam (NL).		
(71) Applicant (for AU BB CA GB IE LK MN MW NZ S. UNILEVER PLC [GB/GB]; Unilever House, Bla London EC4 4BQ (GB).		Published
(71)(72) Applicants and Inventors: HAMERS, R [BE/BE]; Vijversweg 15, B-1640 Sint-Genesius-Ro HAMERS-CASTERMAN, Cécile [BE/BE]; Vij		.

(54) Title: PRODUCTION OF ANTIBODIES OR (FUNCTIONALIZED) FRAGMENTS THEREOF DERIVED FROM HEAVY CHAIN IMMUNOGLOBULINS OF CAMELIDAE

(57) Abstract

A process is provided for the production of an antibody or a fragment or functionalized fragment thereof using a transformed lower eukaryotic host containing an expressible DNA sequence encoding the antibody or (functionalized) fragment thereof, wherein the antibody or (functionalized) fragment thereof is derived from a heavy chain immunoglobulin of Camelidae and is devoid of light chains, and wherein the lower eukaryotic host is a mould, preferably belonging to the genera Aspergillus or Trichoderma, or a yeast, preferably belonging to the yeast genera Saccharomyces, Kluyveromyces, Hansenula, or Pichia. The heavy chain fragment can contain at least the whole variable domain. A complementary determining region (CDR) different from the CDR belonging to the natural antibody ex Camelidae can be grafted on the framework of the variable domain of the heavy chain immunoglobulin. The catalytic antibodies can be raised in Camelidae against transition state molecules. The functionalized antibody or fragment thereof can comprise a fusion protein of both a heavy chain immunoglobulin from Camelidae or a fragment thereof and another polypeptide, e.g., an enzyme, preferably an oxido-reductase. Also provided are new products obtainable by a process as described, and compositions containing a product produced by a process as described, which composition may contain a new product as provided.

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Title: Production of antibodies or (functionalized) fragments thereof derived from heavy chain immunoglobulins of Camelidae

The present invention relates to a process for the production of antibodies or (functionalized) fragments thereof derived from heavy chain immunoglobulins of *Camelidae* and is partly based on research investigations carried out at the Free University of Brussels. A draft publication thereon already submitted to the periodical Nature and communicated to the present applicants by Prof. R. Hamers reads as follows.

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FUNCTIONAL HEAVY CHAIN IMMUNOGLOBULINS IN THE CAMELIDS

Random association of V_L and V_H repertoires contributes considerably to antibody diversity (1). The diversity and the affinity are then increased by hypermutation in B-cells located in germinal centres (2). Except in the heavy chain disease (3), naturally occurring heavy chain antibodies have not been described, although antigen binding has been demonstrated for separated heavy chains (4) or cloned V_H domains (5). The presence of considerable amounts IgG like material of 100 Kd in the serum of the camel (*Camelus dromedarius*) (6) was confirmed. These molecules are composed of heavy chain dimers and are devoid of light chains. Nevertheless they bear an extensive antigen binding repertoire, a finding which questions the role of the light chains in the camel. Camel heavy chain IgGs lack the C_H1, which in one IgG class might be structurally replaced by an extended hinge. Heavy chain IgGs are a feature of all camelids. These findings open perspectives in engineering of antibodies.

By a combination of affinity chromatography on Protein A and Protein G, three quantitatively important fractions corresponding to subclasses of IgG can be isolated from the serum of camels (Camelus dromedarius) (Fig. 1A, lanes c-f).

One fraction (IgG₁) contains molecules of 170 Kd (Fig. 1B, lane 2) which upon reduction yield 50 Kd heavy chains and large 30 kD light chains (Fig. 1C, lane 2). The two other immunoglobulin fractions contain molecules of approximately 100 Kd

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(Fig. 1B, lanes 1 and 3) which upon reduction yield only heavy chains of respectively 46 Kd (IgG₂ fraction binding only to Protein A) (Fig. 1C, lane 3) and 43 Kd (IgG₃ fraction binding to Protein A and Protein G) (Fig. 1C, lane 1). These two IgG classes appear to lack the light chain completely.

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To exclude the possibility that the light chains were only weakly associated with the heavy chains and lost during the selective purification, whole serum was size fractionated by gel filtration. Coomassie blue staining of unreduced fractions revealed the sequential elution of the 170 Kd IgG₁ followed by the incompletely resolved isotypes IgG₂ and IgG₃ (90 Kd) (Fig. 1D, upper inset). Immunostaining of the same fractions after reduction confirmed that the light chains were present solely in the 50 Kd heavy chain containing fractions (Fig. 1D, lower inset).

A comparative study of old world camelids (Cameles bactrianus and Camelus dromedarius) and new world camelids (Lama pacos, Lama glama and Lama vicugna) showed that heavy chain immunoglobulins are abundant in the sera of all species examined (data not shown) and total up to 75% of the molecules binding to protein A.

The abundance of the heavy chain immunoglobulins in the serum of camelids raises the question as to whether they bear an extensive antigen binding repertoire. This question could be answered by examining the IgG₁, IgG₂ and IgG₃ fractions from the serum of camels (Camelus dromedarius) with a high antitrypanosome titer (7). In radio-immunoprecipitation, purified fractions of IgG₁, IgG₂ and IgG₃ derived from infected camels were shown to bind a large number of antigens present in a ³⁵S methionine labelled trypanosome lysate (Fig. 2A), indicating an extensive repertoire complexity for the three IgG classes. Conversely, in blotting experiments, ³⁵S methionine labelled trypanosome lysate binds to SDS-PAGE separated IgG₁, IgG₂ and IgG₃ obtained from infected animals (Fig. 2B). These findings indicate that the heavy chains alone can generate an extensive repertoire and question the obligatory contribution of the light chain to the useful antibody repertoire in the camelids.

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The camelid $\gamma 2$ and $\gamma 3$ chains are considerably shorter than the normal mammalian γ or camel $\gamma 1$ chains. This would suggest that, as in the case of heavy chain disease (3), deletions have occurred in the $C_{H}1$ protein domain (8,9). To address this question, cDNA was synthesized from camel spleen mRNA and the sequences between the 5' end of the V_{H} and the $C_{H}2$ were amplified by a Polymerase Chain Reaction (PCR), and cloned. Seventeen clones presenting a different V_{H} sequence were isolated and sequenced. Their most striking feature was the complete lack of the $C_{H}1$ domain, the last framework (FR4) residues of the V_{H} region being immediately followed by the hinge (Fig. 3, lower part). The absence of the $C_{H}1$ domain clarifies two important dilemmas.

First, immunoglobulin heavy chains are normally not secreted unless the heavy chain chaperoning protein or BIP (10) has been replaced by the L chain (11), or alternatively the C_H1 domain has been deleted (3,8,9). Secondly, isolated heavy chains from mammalian immunoglobulins tend to aggregate, but are only solubilized by light chains (8,12) which bind to the C_H1 and the V_H domains (13).

14 of the 17 clones were characterized by a short hinge sequence with a length equal to that of human IgG_2 and IgG_4 (14) (Fig. 3). The other 3 had a long hinge sequence containing the 'EPK' hinge motif found in human IgG_1 and IgG_3 (14). They possess the C_{H2} 'APELL/P' motif also found in human IgG_1 and IgG_3 (see SEQ. ID. NO: 1-2), and which is associated with mammary transport of bovine IgG_1 (15). On basis of molecular weight, we expect the "short hinge" clones to correspond to IgG_3 and the "long hinge" clones to IgG_2 .

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In the short hinge containing antibody, the extreme distance between the extremities of the V_H regions will be of the order of 80 Å corresponding to twice the size of a single domain of 40 Å $(2xV_H)$ (16). This could be a severe limitation for agglutinating, cross linking or complement fixation (17,18). In the long hinge containing immunoglobulin the absence of C_H1 might be compensated by the extremely long hinge itself, composed of a 12 fold repeat of the sequence Pro-X (X=Gln, Glu, Lys) (Fig. 3 & 4). NMR (19) and molecular modelling (20) of Pro-X repeats present in

the TonB protein of E. coli (X = Glu, Lys) and the membrane procyclin of trypanosomes (X = Asp, Glu) indicate that these repeated sequences function as rigid rodlike spacers with a diameter of 8 Å and a rise of 2.9 Å per residue. Assuming the same geometry, the long hinge would be 70 Å which compensates for the absence of the $C_{11}1$ domain.

The binding site of heavy chain antibodies cannot form the pocket resulting from adjoining light and heavy chain V regions and the residues of the V_H which normally interact with V_L will be exposed to solvent (3,5,13). It was found that leucine at position 45 conserved in 98% of human and murine V_H sequences (14), and crucial in the V_H-V_L association (13), can be replaced by an arginine (Fig. 3, upper part). This substitution is in accordance with both the lost contact with a V_L domain and an increased solubility.

Unlike myeloma heavy chains which result mainly from C_H1 deletion in a single antibody producing cell (21) the camelid heavy chain antibodies have emerged in a normal immunological environment and it is expected that they will have undergone the selective refinement in specificity and affinity accompanying B cell maturation (1, 2). The obtention of camelid heavy chain antibodies could therefore be an invaluable asset in the development and engineering of soluble V_H domains (5) or of new immunologicals for diagnostic, therapeutic or biochemical purposes.

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Figure 1 Characterisation and purification of camel IgG classes on Protein A, Protein G and gel filtration.

- (A) The fraction of *C. dromedarius* serum adsorbed on Protein A shows upon reduction on SDS-PAGE three heavy chain components of respectively 50, 46, and 43 Kd (bands between dots), absent in the non adsorbed fraction (lane d), and light chain components of around 30 Kd (lane c) considerably larger than rabbit light chain (lane a, rabbit IgG). The fractions adsorbed on Protein G (lane e) lack the 46 Kd heavy chain which remains in the non adsorbed fraction (lane f). Lane b contains a size marker.
- 10 (B and C) By differential adsorption and elution on Protein G and Protein A, the IgG fractions containing 43 Kd (lane 1), 46 Kd (lane 3) and 50 Kd (lanes 2) heavy chains were purified and analysed on SDS-PAGE in absence (B) or presence (C) of DTT.
- (D) Whole camel serum (0.1 ml) was fractionated by gel filtration on a Superdex 200 column using 150 mM NaCl, 50 mM sodium phosphate buffer pH 7.0 as eluent. Affinity purified IgG₂ and IgG₃ elute at the positions indicated by arrows. The fractions of interest were further analysed by SDS-PAGE with or without prior reduction. The protein contents as visualized by Coomassie blue (without reduction, upper inset) are compared with the immunoglobulins from the same fractions (after reduction with DTT, lower inset) as revealed by Western blotting with a rabbit anticamel-IgG (lower inset).

METHODS. 5 ml of *C. dromedarius* serum is adsorbed onto a 5 ml Protein G
Sepharose (Pharmacia) column, and washed with 20 mM phosphate buffer, pH 7.0.

25 Upon elution with 0.15 M NaCl, 0.58 % acetic acid (pH 3.5), IgG₃ of 100 Kd is eluted which upon reduction yields heavy chains of 43 Kd (lane 1, B and C). IgG₁ of 170 Kd can subsequently be eluted with pH 2.7 buffer (0.1 M Gly-HCl). This fraction, upon reduction, yields a 50 Kd heavy chain and a broad light chain band (lane 2, C). The fraction not adsorbed on Protein G is brought on a 5 ml Protein A

30 Sepharose column. After washing and elution with 0,15 M NaCl, 0.58% acetic acid (pH 4.5) IgG₂ of 100 Kd is obtained which consists solely of 46 Kd heavy chains (lane 3, C).

- Figure 2 Repertoire complexity and antigen binding capacity of camel IgG₁, IgG₂ and IgG₃ analysed by radioimmunoprecipitation (A) or Western blotting (B & C).
- (A) Serum or purified IgG fractions from healthy or *Trypanoma evansi* infected *C. dromedarius* (CATT titer 1/160 (7)) were incubated with labelled trypanosome lysate, recovered with Protein A Sepharose and analysed by SDS-PAGE. The relative counts recovered are inscribed below each lane. No trypanosome proteins bind to the Protein A or to the healthy camel immunoglobulins.
- 10 (B) 20 μg of IgG₁, IgG₂ and IgG₃ from healthy and trypanosome infected animals were separated by SDS-PAGE without prior reduction or heating. The electroblotted proteins were incubated with the labelled trypanosome lysate. The IgG₂ shows a single antigen binding component corresponding to the heavy chain immunoglobulin whereas the IgG₃ fraction appears to contain in addition two larger antigen binding components barely detectable by Ponceau Red staining (C). These are possibly Ig classes copurified as immunocomplexes present in the serum of the infected animals.
- METHODS. (35S)-methionine labelled *Trypanosoma evansi* lysate (500,000 counts)

 (22) was incubated (4°C, 1 hour) with 10 μl of serum or, 20 μg of IgG₁, IgG₂ or IgG₃ in 200 μl of 0.4 M NaCl, 10 mM EDTA, 10 mM Tris (pH 8.3), containing 0.1 M TLCK. 10 mg of Protein A SeDharose suspended in 200 μl of the same buffer was added (4°C, 1 hour). After washing and centrifugation, each pellet was resuspended in 75 μl SDS PAGE sample solution containing DTT, and heated for 3 min. at 100°C. After centrifugation, 5 μl of the supernatant was saved for radioactivity counting and the remainder analysed by SDS PAGE and fluorography.

 The nitrocellullose filter of the Western blot of purified fractions IgG₁, IgG₂ and IgG₃ was stained with Ponceau Red (C) or incubated with 1% ovalbumin in TST buffer (Tris 10 mM, NaCl 150 mM, Tween 0,05%) (B). The membrane was extensively washed with TST buffer and incubated for 2 hours with (35S)-labelled trypanosome antigen. To avoid unspecific binding, the labelled trypanosome antigen

lysate was filtered (45 μ) and incubated with healthy camel immunoglobulin and ovalbumin adsorbed on a nitrocellulose membrane.

Figure 3 Amino acid sequences of the V_{II} framework, and hinge/ C_{II} 2 of Camelus dromedarius heavy chain immunoglobulins, compared to human (italic) V_{II} framework (subgroup III) and hinges of human IgG (14).

METHODS. Total RNA was isolated from a dromedary spleen (23), mRNA was purified with oligo T-paramagnetic heads (PolyATract-Promega). 1 µg mRNA was used for preparing double-strand cDNA (23) after an oligo-dT priming using 10 enzymes provided by Boehringer Mannheim. 5 µg of cDNA was amplified by PCR in a 100 µl reaction mixture (10mM Tris-HCl pH 8.3, 50 mM KC1,15 mM MgCl₂, 0.01% (w/v) gelatine, 200 µM of each dNTP). 25 pmoles of each oligonucleotide of the mouse V_{II} (24), containing a XhoI site, and 5'-CGCCATCAAGGTACCAGT-TGA-3' (see SEQ. ID. NO: 3) were used as primers. The 3' end primer was deduced from partial sequences corresponding to y chain amino acid 296 to 288 15 (T.Atarhouch, C. Hamers-Casterman, G. Robinson, private communication) in which one mismatch was introduced to create a KDnI restriction site. After a round of denaturing annealing (94°C for 5 min. and 54°C for 5 min.), 2 U of Taq DNA polymerase were added, to the reaction mixture before subjecting it to 35 cycles of amplification (5). The PCR products were purified by phenol-chloroform extraction followed by HPLC (Genpak-fax column, Waters) and finally by MERMAID (BIO 101, Inc.). After these purification steps, the amplified cDNA was digested with XhoI and KpnI, and ligated into pBluescript.

The clones were sequenced by the dideoxy chain termination method (25). The sequences were translated into amino acids which allowed their assignment to well defined domains of the Ig molecule (14); see SEQ. ID. NO: 4-12

Figure 4 Schematic representation of the structural organisation of the camel immunoglobulins (adapted from 26).

On the basis of size consideration, the IgG₁ fraction possess probably the normal antibody assembly of two light and two heavy chains. IgG₃ would have a hinge comparable in size to the human IgG₁, IgG₂ and IgG₄. The two antigen binding sites

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are much closer to each other as this camel IgG lacks the $C_{11}1$ domain. In the camel IgG₂ the long hinge, being formed of Pro-X repeats (X = Glu, Gln or Lys), most likely adopt a rigid structure (19,20). This long hinge could therefore substitute the $C_{11}1$ domain and bring the two antigen binding sites of IgG₂ to normal positions.

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--- End of Draft publication ---

Background of the invention

Already at a very early stage during evolution antibodies have been developed to 10 protect the host organisms against invading molecules or organisms. Most likely one of the earliest forms of antibodies must have been developed in Agnatha. In these primitive fishes antibodies of the IgM type consisting of heavy and lights chains have been detected. Also in many other forms of life ranging from amphibians to mammals antibodies are characterized by the feature that they consist of two heavy and two light chains, although the heavy chains of the various classes of immunoglobulins are quite different. These heavy and light chains interact with each other by a number of different physical forces, but interactions between hydrophobic patches present on both the heavy and light chain are always important. The interaction between heavy and light chains exposes the complementarity determining 20 regions (CDRs) of both chains in such a way that the immunoglobulin can bind the antigen optimally. Although individual heavy or light chains have also the capability to bind antigens (Ward et al., Nature 341 (1989) 544-546 = ref. 5 of the above given draft publication) this binding is in general much less strong than that of combined heavy and light chains.

25 Heavy and light chains are composed of constant and variable domains. In the organisms producing immunoglobulins in their natural state the constant domains are very important for a number of functions, but for many applications of antibodies in industrial processes and products their variable domains are sufficient. Consequently many methods have been described to produce antibody fragments.

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One of these methods is characterized by cleavage of the antibodies with proteolytic enzymes like papain and pepsin resulting in (a) antibody fragment comprising a light

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chain bound via an S-S bridge to part of a corresponding heavy chain formed by proteolytic cleavage of the heavy chain (Fab), or (b) a larger fragment of the antibody comprising two of these Fabs still connected to each other via an S-S bridge in enlargements of the heavy chain parts, indicated with F(ab)₂, respectively (see patent applications EP-A-0125023 (GENENTECH / Cabilly et al., 1984) and WO-A-93/02198 (TECH. RES. CENT. FINLAND / Teeri et al., 1993) for definitions of these abbreviations). The disadvantage of the enzymatic route is that the production of whole antibodies is expensive and the enzymatic processing increases the costs of these fragments even more. The high costs of antibody fragments block the application of these fragments in processes and products outside the pharmaceutical industry.

Another method is based on linkage on DNA level of the genes encoding (parts of) the heavy chain and the light chain. This linkage and the subsequent production of these chimeric immunoglobulins in microorganisms have been described (for Fab fragments see e.g. Better et al., Science 240 (1988) 1041-1043, for F_v fragments (combination of variable fragments of the heavy chain (V_H) and light chain (V_L) still connected to each other by non-covalent binding interactions) see e.g. Skerra et al., Science 240 (1988) 1938, and for single chain F_v fragments (ScF_v; an F_v fragment in which the two variable fragments are linked to each other by a linker peptide) see e.g. Bird et al., Science 242 (1988) 423-426. Provided that an appropriate signal sequence has been placed in front of the single chain V_H and V_L antibody fragment (ScF_v), these products are translocated in E. coli into the periplasmic space and can be isolated and activated using quite elaborate and costly procedures. Moreover the application of antibody fragments produced by E. coli in consumer products requires extensive purification processes to remove pyrogenic factors originating from E. coli. For this and other reasons the production of ScF, in microorganisms that are normally used in the fermentation industry, like prokaryotes as Streptomyces or Bacillus (see e.g. Wu et al. Bio/Technology 11 (1993) 71) or yeasts belonging to the genera Saccharomyces (Teeri et al., 1993, supra), Kluyveromyces, Hansenula, or Pichia or moulds belonging to the genera Aspergillus or Trichoderma is preferred. However with a very few exceptions the production of ScF, antibodies using these systems

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proved to be impossible or quite poor. Although the exact reasons for the poor production are not well known, the use of linkers between the V_{II} and V_{L} chains not designed for secretion (Teeri *et al.*, 1993, *supra*) may be a reason.

Another reason may be incorrect folding of ScF_v. The frameworks and to a limited extend the CDRs of variable domains of light and heavy chains interact with each other. It has been described by Chothia et al. (J. Mol. Biol. 186 (1985) 651-663 = ref. 13 of the above given draft publication) that this interaction involves amino acids at the following positions of the variable region of the heavy chain: 35, 37, 39, 44-45, 47, 100-103 and 105 (numbering according to Kabat et al., In "Sequences of Proteins of Immunological Interest, Public Health Service, NIH, Washington DC, 1983 = ref. 14 of the above given draft publication). Especially leucine at position 45 is strongly conserved and the whole apolar side chain of this amino acid seems to be involved in the interaction with the light chain. These strong interactions may fold the ScF_v into a structure that can not be translocated in certain types of lower eukaryotes.

Thus the use of a linker in the production of ScF_{ν} for connecting a V_H chain to a V_L chain, might negatively influence either the translocation, or the folding of such ScF_{ν} or both.

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Not prior-published European patent application 92402326.0 filed 21.08.92 (C. Casterman & R. Hamers) discloses the isolation of new animal-derived immunoglobulins devoid of light chains (also indicated as heavy chain immunoglobulins), which can especially originate from animals of the camelid family (Camelidae). This European patent specification, now publicly available as EP-A1-0 584 421, is incorporated herein by reference. These heavy chain immunoglobulins are characterized in that they comprise two heavy polypeptide chains sufficient for the formation of one or more complete antigen binding sites, whereby a complete antigen binding site means a site which will alone allow the recognition and complete binding of an antigen, which can be verified by any known method regarding the testing of the binding affinity. The European patent specification further discloses methods for

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isolating these heavy chain immunoglobulins from the serum of *Camelidae* and details of the chemical structure of these heavy chain immunoglobulins. It also indicates that these heavy chain immunoglobulins and derivatives thereof can be made by using recombinant DNA technology in both prokaryotes and eukaryotes. The present invention relates to a further development of the work disclosed in that prior-filed but not prior-published European specification.

Due to the absence of light chains in most of the immunoglobulins of *Camelidae* such linkers are not necessary, thereby avoiding the above-mentioned potential problems.

As described above in the draft publication for Nature, now publicly available as Nature 363 (3 June 1993) 446-448, and in the not prior-published European patent application 92402326.0 (supra) it was surprisingly found that the majority of the protein A-binding immunoglobulins of Camelidae consists just of two heavy chains and that these heavy chains are quite different from common forms of heavy chains, as the C_H1 domain is replaced by a long or short hinge (indicated for IgG₂ and IgG₃, respectively, in Figure 4 of the above given draft publication for Nature). Moreover these heavy chains have a number of other features that make them remarkably different from the heavy chains of common immunoglobulins.

One of the most significant features is that they contain quite different amino acid residues at those positions involved in binding to the light chain, which amino acids are highly conserved in common immunoglobulins consisting of two heavy and two light chains (see Table 1 and SEQ. ID. NO: 13-31).

Table 1 Comparison af amino acid sequences of various immunoglobulins Alignment of a number of V_{II} regions of Camel heavy chain antibodies compared with those of mouse (M, top line) and human (H, second line). Framework fragments are indicated in capitals, CDR fragments in small print; see SEQ. ID. NO: 13-31 for sequences indicated by M, H, 1, 2, 3, 7, 9, 11, 13, 16, 17, 18, 19, 20, 21, 24, 25, 27, 29, respectively.

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          EVKLVESGGG LVQPGGSLRL SCATSGFTFS dfyme..WVR QPPGKRLEWI
       m
          EVOLVESGGG LVOPGGSLRL SCAASGFTFS syams..WVR QAPGKGLEWV
       h
    cam1
           .....GG SVQAGGSLRL SCAASGYSNC pltws..WYR QFPGTEREFV
    cam2
          DVQLVASGGG SVQAGGSLRL SCTASGDSFS rfams..WFR QAPGKECELV
    cam3
           ......GG SVQTGGSLRL SCAVSGFSFS tscma..WFR QASGKQREGV
15
    cam7
           ......GG SVQGGGSLRL SCAISGYTYG sfcmg..WFR EGPGKEREGI
    cam9
           ......GG SVQAGGSLTL SCVYTNDTGT ...mg..WFR QAPGKECERV
   cam11
           ......GG SVQAGGSLRL SCNVSGSPSS tyclg..WFR QAPGREREGV
   cam13
           ......GG SVEAGGSLRL SCTASGYVSS ...ma..WFR QVPGQEREGV
           ......GG SAQAGGSLRL SCAAHGIPLN gyyia..WFR QAPGKGREGV
   cam16
20
   cam17
           .....GG SVQPGGSLTL SCTVSGATYS dysig..WIR QAPGKDREVV
   cam18
           ......GG SVQAGGSLRL SCTGSGFPYS tfclg..WFR QAPGKEREGV
   cam19
           ......GG SVQAGGSLRL SCAASDYTIT dycma..WFR QAPGKERELV
   cam20
           ......GG SVQVGGSLRL SCVASTHTDS stcig..WFR QAPGKEREGV
   cam21
           ......GG SVQVGGSLKL SCKISGGTPD rvpkslaWFR QAPEKEREGI
25
           ......GG SVQAGGSLRL SCNVSGSPSS tyclg..WFR QAPGKEREGV
   cam24
           ......GG SVQTGGSLRL SCEISGLTFD dsdvg..WYR QAPGDECKLV
   cam25
   cam27
           ......GG SVQAGGSLRL SCASSSKYMP ctydmt.WYR QAPGKEREFV
   cam29
           .....exxGG SVQAGGSLRL SCVASGFNFE tsrma..WYR QTPGNVCELV
30
          A..asrnkan dytteysasv kgRFIVSRDT SQSILYLQMN ALRAEDTAIY
       \mathbf{m}
          S..xisxktd ggxtyyadsv kgRFTISRDN SKNTLYLQMN SLRAEDTAVY
          S..smd...p dgntkytysv kgRFTMSRGS TEYTVFLQMD NLKPEDTAMY
    cam1
35
    cam2
          S..siq...s ngrtteadsv qgRFTISRDN SRNTVYLQMN SLKPEDTAVY
          Aainsgggrt yyntyvaesv kgRFAISQDN AKTTVYLDMN NLTPEDTATY
    cam3
    cam7
          A..tiln..g gtntyyadsv kgRFTISQDS TLKTMYLLMN NLKPEDTGTY
    cam9
          A..hit...p dgmtfidepv kgRFTISRDN AQKTLSLRMN SLRPEDTAVY
   cam11
          T..aint..d gsiiyaadsv kgRFTISQDT AKETVHLQMN NLQPEDTATY
40
   cam13
          A..fvqt..a dnsalygdsv kgRFTISHDN AKNTLYLQMR NLQPDDTGVY
   cam16
          A..ting..g rdvtyyadsv tgRFTISRDS PKNTVYLQMN SLKPEDTAIY
          A..aant..g atskfyvdfv kgRFTISQDN AKNTVYLQMS FLKPEDTAIY
   cam17
   cam18
          A..gins..a ggntyyadav kgRFTISQGN AKNTVFLQMD NLKPEDTAIY
          A.aiqvvrsd trltdyadsv kgRFTISQGN TKNTVNLQMN SLTPEDTAIY
   cam19
          A..siyf..g dggtnyrdsv kgRFTISQLN AQNTVYLQMN SLKPEDSAMY
   cam20
          A..vlst..k dgktfyadsv kgRFTIFLDN DKTTFSLQLD RLNPEDTADY
   cam21
          T..aint..d gsviyaadsv kgRFTISQDT AKKTVYLQMN NLQPEDTATY
   cam24
          Sgilsdgtpy tksgdyaesv rgRVTISRDN AKNMIYLQMN DLKPEDTAMY
   cam25
          S..sin...i dgkttyadsv kgRFTISQDS AKNTVYLQMN SLKPEDTAMY
   cam27
50
   cam29
          S..siy...s dgktyyvdrm kgRFTISREN AKNTLYLQLS GLKPEDTAMY
```

Table 1 (Cont.) Comparison af amino acid sequences of various immunoglobulins Alignment of a number of V_{II} regions of Camel heavy chain antibodies compared with those of mouse (M, top line) and human (H, second line). Framework fragments are indicated in capitals, CDR fragments in small print; see SEQ. ID. NO: 13-31 for sequences indicated by M, H, 1, 2, 3, 7, 9, 11, 13, 16, 17, 18, 19, 20, 21, 24, 25, 27, 29, respectively.

```
101
10
           YCARdyygss .....y.. f.....dvWG AGTTVTVSS
        m
        h
           YCARxxxxx xxxxxyyyyh x....fdyWG QGTLVTVSS
           YCKTalqpgg ycgygx.....clWG QGTQVTVSS
YCGAvslmdr isqh.....gcRG QGTQVTVSL
     caml
     cam2
     cam3
           YCAAvpahlg pgaildlkky .....kyWG QGTQVTVSS
           YCAAelsggs celpllf......dyWG QGTQVTVSS YCAAdwkywt cgaqtggyf.....gqWG QGAQVTVSS
15
     cam7
     cam9
    camll
           YCAArltemg acdarwatla trtfaynyWG QGTQVTVSS
           YCAAqkkdrt rwaeprew.....nnWG QGTQVTASS
FCAAgsrfss pygstsrles .sdy..nyWG QGIQVTASS
    cam13
    cam16
20
           YCAAadpsiy ysilxiey.....kyWG QGTQVTVSS
    cam17
           YCAAdspcym ptmpappird sfgw..ddFG QGTQVTVSS
    cam18
    cam19
           SCAAtssfyw ycttapy.....nvWG QGTQVTVSS
           YCAIteiewy gcnlrttf......trWG QGTQVTVSS
    cam20
           YCAAnqlagg wyldpnywls vgay..aiWG QGTHVTVSS
    cam21
25
    cam24
           YCAArltemg acdarwatla trtfaynyWG RGTQVTVSS
    cam25
           YCAVdgwtrk eggiglpwsv qcedgynyWG QGTQVTVSS
    cam27
           YCKIdsypch 11......dvWG QGTQVTVSS
           YCAPveypia dmcs.....ryGD PGTQVTVSS
    cam29
30
```

For example, according to Pessi et al. (1993) a subdomain portion of a V_H region of common antibodies (containing both heavy chains and light chains) is sufficient to direct its folding, provided that a cognate V_L moiety is present. Thus it might be expected from literature on the common antibodies that without V_L chains proper folding of heavy chains cannot be achieved. A striking difference between the common antibodies and the Camelidae-derived heavy chain antibodies is, that the highly conserved apolar amino acid leucine (L) at place 45 present in common antibodies is replaced in most of the Camelidae-derived heavy chain antibodies by the charged amino acid arginine (R), thereby preventing binding of the variable region of the heavy chain to that of the light chains.

Another remarkable feature is that one of the CDRs of the heavy chains of this type

of immunoglobulins from Camelidae, CDR3, is often much longer than the

corresponding CDR3 of common heavy chains. Besides the two conserved cysteines forming a disulphide bridge in common V_H fragments, the *Camelidae* V_H fragments often contain two additional cysteine residues, one of which often is present in CDR3.

According to the present inventors these features indicate that CDR3 may play an important role in the binding of antigens by these heavy chain antibodies and can compensate for the absence of light chains (also containing CDRs) in binding of antigens by immunoglobulins in *Camelidae*.

Thus, as the heavy chains of *Camelidae* do not have special features for interacting with corresponding light chains (which are absent), these heavy chains are very different from common heavy chains of immunoglobulins and seem intrinsically more suitable for secretion by prokaryotic and lower eukaryotic cells.

The present inventors realized that these features make both intact heavy chain immunoglobulins of *Camelidae* and fragments thereof very attractive for their production by microorganisms. The same holds for derivatives thereof including functionalized fragments. In this specification the term "functionalized fragment" is used for indicating an antibody or fragment thereof to which one or more functional groups, including enzymes and other binding polypeptides, are attached resulting in fusion products of such antibody fragment with another biofunctional molecule.

Summary of the invention

In a broad sense the invention provides a process for the production of an antibody or a fragment or functionalized fragment thereof using a transformed lower eukaryotic host containing an expressible DNA sequence encoding the antibody or (functionalized) fragment thereof, wherein the antibody or (functionalized) fragment thereof is derived from a heavy chain immunoglobulin of Camelidae and is devoid of light chains, and wherein the lower eukaryotic host is a mould or a yeast. Thus the lower eukaryotic host can be a mould, e.g. belonging to the genera Aspergillus or Trichoderma, or a yeast, preferably belonging to the yeast genera Saccharomyces, Kluyveromcyes, Hansenula, or Pichia. Preferably the fragments still contain the whole variable domain of these heavy chains.

The invention also provides methods to produce such heavy chain immunoglobulins or (functionalized) fragments thereof in which methods the framework or the CDRs of these heavy chains are modified by random or directed mutagenesis in such a way that the mutated heavy chain is optimized for secretion by the host microorganism into the fermentation medium.

Another embodiment of the invention is that CDRs can be grafted on these optimized frameworks (compare grafting of CDRs on human immunoglobulins as described by e.g. Jones et al., Nature 321 (1986) 522). These CDRs can be obtained from common antibodies or they may originate from heavy chain immunoglobulins of Camelidae. The binding properties may be optimized by random or directed

mutagenesis. Thus in a process according to the invention an antibody or (functionalized) fragment thereof derived from a heavy chain immunoglobulin of Camelidae can be produced which comprises a CDR different from the CDR belonging to the natural antibody ex Camelidae which is grafted on the framework of the variable domain of the heavy chain immunoglobulin ex Camelidae.

The invention also provides a method for the microbiological production of catalytic antibodies. These antibodies are preferably raised in *Camelidae* against transition state molecules following procedures similar to the one described by Lerner *et al.*, Science 252 (1991) 659-667. Using random or site-directed mutagenesis such

catalytic antibodies or fragments thereof can be modified in such a way that the catalytic activity of these (functionalized) antibodies or fragments can be further improved.

For preparing modified heavy chain antibodies a process according to the invention is provided, in which the DNA sequence encodes a modified heavy chain immunoglobulin or a (functionalized) fragment thereof derived from *Camelidae* and being devoid of light chains, and is made by random or directed mutagenesis or both. Thus the resulting immunoglobulin or (functionalized) fragment thereof is modified such that

- it is better adapted for production by the host cell, or
- it is optimized for secretion by the lower eukaryotic host into the fermentation medium, or
 - its binding properties (k_{on} and k_{off}) are optimized, or

- its catalytic activity is improved, or
- it has acquired a metal chelating activity, or
- its physical stability is improved.
- Another particular embodiment of the present invention relates to genes encoding fusion proteins consisting of both a heavy chain immunoglobulin from Camelidae or part thereof and a second protein or another polypeptide, e.g. an enzyme, in particular an oxido-reductase, and to expression products of such genes. By means of the heavy chain immunoglobulin (fragment) the protein or enzyme can be guided to a target thereby increasing the local efficiency of the protein or enzyme significantly. Thus according to this embodiment of the invention a process is provided, in which the functionalized antibody or fragment thereof comprises a fusion protein of both a heavy chain immunoglobulin from Camelidae or a fragment thereof and another polypeptide, e.g. an enzyme, preferably an oxido-reductase.

- As a result of a process according to the invention known products may be produced, e.g. antibodies also produced by *Camelidae*, but many of the possible products will be new products, thus the invention also provides new products obtainable by a process according to the invention.
- The products so produced can be used in compositions for various applications.

 Therefore, the invention also relates to compositions containing a product produced by a process according to the invention. This holds for both old products and new products.

25 Brief Description of the Figures

Figures 1-4 were already described above in the draft publication.

Figure 1 Characterisation and purification of camel IgG classes on Protein A, Protein G and gel filtration.

Figure 2 Repertoire complexity and antigen binding capacity of camel IgG₁,

IgG₂ and IgG₃ analysed by radioimmunoprecipitation (A) or

Western blotting (B & C).

	Figure 3	mino acid sequences of the V_H framework, and hinge/ C_{i1} 2 of	
		Camelus dromedarius heavy chain immunoglobulins, compared to	
		human (italic) V _{II} framework (subgroup III) and hinges of human	
		IgG (14); see SEQ. ID. NO: 4-12.	
5	Figure 4	Schematic representation of the structural organisation of the camel	
		immunoglobulins (adapted from 26).	
	Figure 5	DNA and amino acid sequences of the Camel V _{II} fragments fol-	
		lowed by the Flag sequence as present in pB03 (Figure 5A), pB09	
		(Figure 5B) and pB24 (Figure 5C); see SEQ. ID. NO: 32-37.	
10	Figure 6	Nucleotide sequence of synthetic DNA fragment cloned into	
		pEMBL9 (Example 1); see SEQ. ID. NO: 38-41.	
	Figure 7	Schematic drawing of plasmid pUR4423	
	Figure 8	Schematic drawing of plasmid pUR4426	
	Figure 9	Schematic drawing of plasmid pUR2778	
15	Figure 10	Schematic drawing of plasmid pUR4429	
	Figure 11	Schematic drawing of plasmid pUR4430	
	Figure 12	Schematic drawing of plasmid pUR4445	
	Figure 13	Schematic drawing of plasmid pUR4446	
	Figure 14	Schematic drawing of plasmid pUR4447	
20	Figure 15	Schematic drawing of plasmid pUR4451	
	Figure 16	Schematic drawing of plasmid pUR4453	
	Figure 17	Schematic drawings of plasmids pUR4437 and pUR4438	
	Figure 18	Schematic drawings of plasmids pUR4439 and pUR4440	
	Figure 19	Nucleotide sequence of synthetic DNA fragment cloned into	
25		pEMBL9 (Example 6); see SEQ. ID. NO: 42-45.	
	Figure 20	Schematic drawing of plasmid pAW14B.	
	Figure 21	Western blot analysis of culture medium of S. cerevisiae trans-	
		formants containing pUR4423M (see A) or pUR4425M (see B).	
		Samples were taken after 24 (see 1) or 48 hours (see 2). For	
30		pUR4425M two bands were found due to glycosylation of the	
		antibody fragment.	

Detailed description of the invention

The present invention relates to the production of antibodies or (functionalized) fragments thereof derived from heavy chain immunoglobulins of *Carnelidae* by eukaryotes, more in particular by lower eukaryotes such as yeasts and fungi.

Therefore, mRNA encoding immunoglobulins of Camelidae was isolated and transcribed into cDNA according to the procedures described in the above given draft publication and not prior-published European patent application 92402326.0. In each case primers for the PCR reaction directed to the N-terminus of the V_H domain and PCR primers that either hybridize with the C-terminal regions of the V_H domain or with the short or large hinge regions as described in the above given draft publication, or with the C-terminal region of the C_H2 or C_H3 domains can be used. In this way structural genes can be obtained encoding the following fragments of heavy chain immunoglobulins of Camelidae (Table 2).

15

Table 2. The various forms of immunoglobulins of Camelidae that can be expressed in microorganisms.

- a. the variable domain of a heavy chain;
- 20 b. the variable domain and the short hinge of a heavy chain;
 - c. the variable domain and the long hinge of a heavy chain;
 - d. the variable domain, the C_H2 domain, and either the short or long hinge of a heavy chain;
 - e. a complete heavy chain, including either the short or long hinge.

25

According to procedures described in detail in the Examples these cDNAs can be integrated into expression vectors.

Known expression vectors for Saccharomyces, Kluyveromcyes, Hansenula, Pichia and Aspergillus can be used for incorporating a cDNA or a recombinant DNA according to the invention. The resulting vectors contain the following sequences that are required for expression: (a) a constitutive, or preferably an inducible, promoter; (b) a leader or signal sequence; (c) one of the structural genes as described in Table 2

and (d) a terminator. If the vector is an episomal vector, it preferably comprises an origin of replication as well as a selection marker, preferably a food grade selection marker, (EP-A-487159, UNILEVER / Leenhouts et al.). If the vector is an integration vector, then it preferably comprises sequences that ensure integration and a selection marker in addition to the sequences required for expression of the structural gene encoding a form of the heavy chain immunoglobulin of Camelidae or derivatives thereof. The preferred sequences for integration are sequences encoding ribosomal DNA (WO 91/00920, 1991, UNILEVER / Giuseppin et al.) whereas the selection marker will be preferably a food grade marker.

For Saccharomyces the preferred inducible promoter is the GAL7 promoter (EP-A-0255153, UNILEVER / Fellinger et al.); for Kluyveromyces the preferred inducible promoter is the inulinase promoter (not yet published EP application 92203932.6, UNILEVER / Toschka & Verbakel, which is incorporated herein by reference); for Hansenula or Pichia the preferred inducible promoter is the methanol-oxidase promoter (Sierkstra et al., Current Genetics 19 (1991) 81-87) and for Aspergillus the preferred inducible promoter is the endo-xylanase promoter (not prior-published PCT application PCT/EP 92/02896, UNILEVER / Gouka et al., now publicly available as WO-A-93/12237, which is incorporated herein by reference).

To achieve efficient secretion of the heavy chain immunoglobulin or parts thereof the leader (secretion) sequences of the following proteins are preferred invertees.

the leader (secretion) sequences of the following proteins are preferred: invertase and α -factor for Saccharomyces, inulinase for Kluyveromyces, invertase for Hansenula or Pichia (Sierkstra et al., 1991 supra) and either glucoamylase or xylanase for Aspergillus (not prior-published PCT application WO-A-93/12237, supra). As food-grade selection markers, genes encoding anabolic functions like the leucine2 and tryptophan3 are preferred (Giuseppin et al. 1991, supra). The present invention describes the heterologous production of (functionalized) derivatives or fragments of immunoglobulins in a microorganism, which immunoglobulins in nature occur not as a composite of heavy chains and light chains, but only as a composite of heavy chains. Although the secretion mechanism of mammals and microorganisms is quite

30 similar, in details there are differences that are important for developing industrial processes.

To obtain frameworks of the heavy chain immunoglobulins, that are optimally secreted by lower eukaryotes, genes encoding several different heavy chains can be cloned into the coat protein of bacteriophages and subsequently the frameworks of these heavy chain immunoglobulins can be mutated using known PCR technology,

e.g. Zhou et al., (1991). Subsequently the mutated genes can be been cloned in Saccharomyces and Aspergillus and the secretion of the mutated genes can be compared with the wild type genes. In this way frameworks optimized for secretion may be selected.

Alternatively these structural genes can be linked to the cell wall anchoring part of cell wall proteins, preferably GPI-linked cell wall proteins of lower eukaryotes, which result in the expression of a chimeric protein on the cell wall of these lower eukaryotes (not prior-published EP application 92202080.5, UNILEVER / Klis et al., now publicly available as International (PCT) patent application WO-A-94/01567, which is incorporated herein by reference).

Both methods have the advantage that the binding parts of the immunoglobulins are well exposed to the surrounding of the cell, microorganism, or phage and therefore can bind antigens optimally. By changing the external conditions the binding rates and dissociation rates of this binding reaction can be influenced. Therefore, these systems are very suitable to select for mutated immunoglobulins that have different binding properties. The mutation of the immunoglobulins can either be obtained by random mutagenesis, or directed mutagenesis based on extensive molecular modelling and molecular dynamical studies.

mRNAs encoding heavy chains of immunoglobulins raised in Camelidae against transition state molecules (Lerner et al., 1991 supra) can be obtained using standard techniques. The structural genes encoding various forms of immunoglobulins according to the invention as summarized in Table 2 can be cloned into the coat protein of bacteriophages or as fusion with the anchoring part of cell wall proteins and can be tested on the catalytic property. In this way immunoglobulins or parts thereof having catalytic properties can be determined and selected. Genes encoding these selected immunoglobulins or parts thereof can be mutated as described before and recloned in bacteriophages, but preferably cloned as chimeric cell wall bound catalysts in lower eukaryotes. By performing appropriate catalytic assays, catalytic

immunoglobulins or parts thereof with improved catalytic properties can be determined and selected using standard techniques.

An important application of antibodies, especially outside the pharmaceutical industry, will be chimeric proteins consisting of the binding part of antibodies and enzymes. In this way catalytic biomolecules can be designed that have two binding properties, one of the enzyme and the other of the antibody. This can result in enzymes that have superior activity. This can be illustrated with the following examples:

- a. If the substrate of the enzymic reaction is produced by an organism or an enzyme is recognized by the binding domain of the antibody, the local concentration of the substrate will be much higher than for enzymes lacking this binding domain and consequently the enzymic reaction will be improved. In fact this is a mimic of vectorial metabolism in cells (compare e.g. Mitchell, (1979) Science 206 1148-1159);
- b. If the substrate of the enzymic reaction is converted into a molecule that kills organisms, then the efficiency and specificity of killing can be increased significantly if the enzyme is equipped with an antibody binding domain that recognizes the target organism (e.g. compare Takahashi et al., (1993) Science 259 1460-1463);

20

The invention will be illustrated by the following Examples without being limited thereto. In previously filed Unilever patent specifications several expression vectors were described, e.g. for the yeasts S. cerevisiae, Kluyveromyces, and Hansenula, and the mould Aspergillus. Examples of these publications are EP-A-0173378

(UNILEVER / Ledeboer et al.), EP-A-0255153, supra, and PCT applications
 WO-A-91/19782 (UNILEVER / van Gorcom et al.) and (not prior-published)
 WO-A-93/12237, supra. The genes encoding antibodies or (functionalized) fragments thereof according to the invention can be incorporated into the earlier described expression vectors or derivatives thereof using procedures well known to a skilled
 person in the art. All techniques used for the manipulation and analysis of nucleic acid materials were performed essentially as described in Sambrook et al. (1989)

(see also ref. 23 of the above given draft publication), except where indicated otherwise.

In the description of the Examples the following endonuclease restriction sites are used:

5	AflII	CITTAAG	Mlu1	AICGCGT
	BspHI	TICATGA	Ncol	CICATGG
	BspHI	TICATGA	Not	GCIGGCCGC
	BstEII	GIGTNACC	Nrul	TCGICGA
	Eagl	CIGGCCG	Sall	GITCGAC
10	<i>Eco</i> RI	GIAATTC	Xhol	CITCGAG
	<i>Hin</i> dIII	AJAGCTT	B bs1	GAAGAC(N) ₂ 1 CTTCTG(N') ₆ 1

Example 1 Construction of cassettes encoding V_{II} fragments originating from Camelidae.

For the production of V_H fragments originating from *Camelidae*, the antibody gene fragments were isolated and cloned as described above in the draft publication. The thus obtained gene fragments encode the V_H region, a short or a long hinge region and about 14 amino acids of the C_H2 region. By using standard molecular biological techniques (e.g. PCR technology), the V_H gene fragments could be subcloned and equipped at their 5'-ends with a gene fragment encoding the *pelB* signal sequence and at their 3'-ends with a gene fragment encoding the Flag tail (13 amino acids). Three of these clones were named pB3, pB9 and pB24 and were deposited at the Centraal Bureau voor Schimmelcultures, Baarn on 20 April 1993 with deposition numbers: CBS 270.93, CBS 271.93 and CBS 272.93, respectively. The DNA and amino acid sequences of the *Camelidae*-V_H fragments followed by the Flag sequence are presented in Figure 5(A-C); see SEQ. ID. NO: 32-37.

1.1 Construction of pUR4421

For the construction of yeast expression plasmids encoding the V_H fragments preceded by the invertase (=SUC2) signal sequence, the α -mating factor prepro-

sequence, or the inulinase signal sequence and followed by either nothing, or a Myc tail or Flag tail, the constructs described below can be prepared.

The multiple cloning site of plasmid pEMBL9 (Denthe et al., 1983) (ranging from the EcoRI to the HindIII site) was replaced by a synthetic DNA fragment having the nucleotide sequence as indicated in Figure 6; see SEQ. ID. NO: 38-41. The 5'-part of this nucleotide sequence comprises an EagI site, the first 4 codons of the Camelidae V_{II} gene fragment and a XhoI site coinciding with codons 5 and 6. The 3'-part comprises the last 5 codons of the Camelidae V_{II} gene (encoding VTVSS; see SEQ. ID. NO: 47) part of which coincides partially with a BstEII site), eleven codons of the Myc tail, and an EcoRI site. The EcoRI site, originally present in pEMBL9, is not functional any more, because the 5'- end of the nucleotide sequence contains AATTT instead of AATTC, indicated in Figure 6 as "(EcoRI)". The resulting plasmid is called pUR4421.

15 1.2 Constructs with Flag tail.

After digesting the plasmid pB3 with XhoI and EcoRI, a DNA fragment of approximately 425 bp was isolated from agarose gel. This fragment codes for a truncated V_H-Flag fragment, missing the first 5 amino acids of the Camelidae V_H. The obtained fragment can be cloned into pUR4421. To this end plasmid pUR4421 can be digested with XhoI and EcoRI, after which the about 4 kb vector fragment can be isolated from an agarose gel. Ligation with the about 425 bp fragment will result in plasmid pUR4421-03F.

1.3 Constructs with Myc tail.

- After digesting the plasmid pB3 with XhoI and BstEII, a DNA fragment of approximately 365 bp was isolated from agarose gel. This fragment codes for a truncated V_{II} fragment, missing both the first 4 (QVKL; see SEQ. ID. NO: 46) and the last 5 (VTVSS; see SEQ. ID. NO: 47) amino acids of the Camelidae V_H fragment.
- The obtained fragment was cloned into pUR4421. To this end plasmid pUR4421 was digested with Xhol and BstEII, after which the about 4 kb vector fragment was isolated from an agarose gel. Ligation with the about 365 bp fragment resulted in

plasmid pUR4421-03M, in which the gene encoding the V₁₁ fragment is reconstituted.

1.4 Constructs encoding V_{11} only.

Upon digesting pUR4421-03M or pUR4421-03F with *BstEil* and *Hindlii*, the vector fragments of about 4.4 kb can be isolated from agarose gel and religated in the presence of a synthetic linker peptide having the following sequence:

BstEII HindIII
GTCACCGTCTCCTCATAATGA
GCAGAGGAGTATTACTTCGA

(see SEQ. ID. NO: 48-49).

In the thus obtained plasmid, pUR4421-03, the Myc tail or Flag tail sequences are removed and the V_H gene fragment is directly followed by a stop codon.

1.5 Other constructs.

- After isolating the gene fragments encoding V_{II}-hinge-C_{II}2 fragments as described above in the draft publication, or encoding the intact heavy chain immunoglobulin, it is possible, e.g. by using PCR technology, to introduce an appropriate restriction enzyme recognition site (e.g. *Eco*Rl or *HindIII*) downstream of the hinge region, downstream of the C_{II}2 region, or downstream of the total gene. Upon isolating a
- 20 XhoI-EcoRI or XhoI-HindIII fragment encoding the V_H fragment with a C-terminal extension, the fragment can be cloned into pUR4421 digested with the same restriction enzymes.
 - In analogy with the construction of pUR4421-03, a number of other constructs can be produced encoding functionalized heavy chain fragments in which a second polypeptide is fused to the C-terminal part of the V_H fragment. Optionally, the V_H fragment and the second polypeptide, e.g. an enzyme, might be connected to each other by a peptide linker.

To this end either the BstEII-HindIII fragment or the BstEII-EcoRI fragment of either pUR4421-03F or pUR4421-03M has to be replaced by another BstEII-HindIII or BstEII-EcoRI fragment. The latter new fragment should code for the last amino acids (VTVSS, see SEO.ID. NO: 47) of the V₁₁ fragment, optionally for a linker peptide, and for the polypeptide of interest e.g. an enzyme. Obviously, the introduction of the DNA fragment should result in an in frame fusion between the

 V_{II} gene fragment and the other DNA sequence encoding the polypeptide of interest.

Alternatively, it is possible to replace the Eagl-XhoI fragment of pUR4421-03 with another DNA fragment, coding for a polypeptide of interest, optionally for a peptide linker, and for the first 4 (QVKL, see SEQ.ID. NO: 46) amino acids of the V_H fragment, resulting in an in frame fusion with the remaining part of the V_H fragment. In this way, it is possible to construct genes encoding functionalized V_H fragments in which the second polypeptide is fused at the N-terminal part of the V_H fragment, optionally via a peptide linker.

Obviously, it is also possible to construct genes encoding functionalized V_H fragments having a polypeptide fused to the N-terminal as well as fused to the C-terminal end, by combining the above described construction routes.

The polypeptides used to functionalize the V₁₁ fragments might be small, like the

Myc and the Flag tails, or intact enzymes, like glucose oxidase, or both.

From all the above described constructs, derived from pUR4421, an appropriate Eagl-HindIII fragment, encoding the functionalized V_H fragment, can be isolated and cloned into a number of different expression plasmids. Several are exemplified in more detail in the following Examples. Although only the V_H fragments are exemplified, similar constructs can be prepared for the production of larger heavy chain fragments (e.g. V_{II}-hinge or V_H-hinge-C_H2) or intact heavy chains. The Eagl site is introduced before the first codon of the V_H fragment, facilitating an in frame fusion with different yeast signal sequences.

In particular cases, were additional Eagl and/or HindIII sites are present in the cloned fragments, it is necessary to perform partial digestions with one or both restriction enzymes.

Although the above and following constructions only consider the V_{II} fragment cloned in pB3, a comparable construction route can be used for the construction of expression plasmids for the production of V_{II} fragments like V_H-09 and V_H-24, or other V_{II} fragments.

Example 2 Construction of S. cerevisiae episomal expression plasmids for Camelidae V_{11} .

For the secretion of recombinant protein from S. cerevisiae it is worthwhile to test in parallel the two most frequently applied homologous signal sequences, the SUC2

invertase signal sequence and the prepro-α mating factor sequence.

The episomal plasmid pSY1 and pSY16 (Harmsen et al., 1993) contain expression

cassettes for the α-galactosidase gene. Both plasmids contain the GAL7 promoter and PGK terminator sequences. pSY1 contains the invertase (SUC2) signal sequence and pSY16 contains a slightly modified (Harmsen et al., 1993) prepro-α-

10 mating factor signal sequence.

Both plasmids, pSY1 and pSY16 can be digested with EagI and HindIII, the about 6500 bp long vector backbone of both plasmids can be isolated and subsequently ligated with the EagI/HindIII fragments from pUR4421-03F (~465 bp), pUR4421-03M (~455 bp) or pUR4421-03 (~405 bp) (See above).

This results in a series of 6 different episomal plasmids for expression in S. cerevisiae, containing behind the SUC2- and the α mating factor prepro-sequence the V_H-Flag coding sequence (designated pUR4423F and pUR4426F), the V_H-Myc coding sequence (designated pUR4423M and pUR4426M) or the coding sequence of V_H followed by a stop codon (designated pUR4423, Figure 7 and pUR4426,

20 Figure 8).

Obviously, it is possible to use promoter systems different from the inducible GAL7 promoter, e.g. the constitutive GAPDH promoter.

2.1 Production of V_{II} -03-myc and V_{II} -24-myc.

25 After introducing the expression plasmids pUR4423M (coding for V_{II}-03-myc, preceded by the SUC2-signal sequence) and pUR4425M (coding for V_{II}-24-myc, preceded by the SUC2-signal sequence) into S. cerevisiae via electroporation, transformants were selected from minimal medium agar plates (comprising 0.7 % yeast nitrogen base, 2 % glucose and 2 % agar, supplemented with the essential amino acids and bases).

For the production of antibody fragments the transformants were grown overnight in selective minimal medium (comprising 0.7 % yeast nitrogen base, 2 % glucose,

supplemented with the essential amino acids and bases) and subsequently diluted ten times in YPGal medium (comprising 1 % yeast extract, 2 % bacto pepton and 5 % galactose). After 24 and 48 hours of growth, samples were taken for Western blot analysis (Figure 21). For the immuno detection of the produced V_{II}-myc fragments monoclonal anti-myc antibodies were used.

In essentially the same way comparable results were obtained with a yeast transformed with pUR4424M containing a DNA sequence encoding the V_{II}-09-myc protein.

Example 3 Construction of *S. cerevisiae* multicopy integration vectors for the expression of *Camelidae* V_{II}.

To combine the benefits of high copy number and mitotically stable expression, the concept of a multicopy integration system into the rDNA locus of lower eukaryotes has already been successfully proven (Giuseppin et al. supra).

One of these vectors is pUR2778, a derivative of pUR2774 (Giuseppin et al. supra) from which the pol1-S.O. reporter gene sequence was removed (Figure 9).

This integrating plasmid, pUR2778, can be used for integration of Camelidae V_H coding sequences, hence the vector can be digested with Sacl and HindIII after

From the in example 2 described pUR4423 or pUR4426 types of plasmids, SacIHindIII fragments can be isolated encoding a V_H fragment preceded by a signal sequence (SUC2 or α mating factor prepro) and followed by nothing or a Myc or Flag tail.

Ligation of these Sacl-HindIII fragments with the 7.3 kb vector fragment will result in integration plasmids, encoding the (functionalized) V_H fragments under the regulation of the strong and inducible GAL7 promoter.

In this way the following expression plasmids were obtained:

which the 7.3 kb vector fragment can be isolated.

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For schematic drawings see Figure 10 for pUR4429 and Figure 11 for pUR4430. Obviously, comparable constructs can be prepared for other heavy chain antibodies or fragments thereof.

As mentioned before, different promoters might be used, for example, the constitutive GAPDH promoter.

Example 4 Construction of expression plasmids for the production of (functionalized) V_{11} fragments from Camelidae by Kluyveromyces

4.1. Construction of Kluyveromyces lactis episomal expression plasmids Camelidae.

Yeast strains of the genus *Kluyveromyces* have been used for the production of enzymes, such as \(\textit{B}\)-galactosidase for many years, and the growth of the strains has been extensively studied. *Kluyveromyces lactis* is well known for the ability to utilize a large variety of compounds as carbon and energy sources for growth. Since these strains are able to grow at high temperatures and exhibit high growth rates, they are promising hosts for industrial production of heterologous proteins (Hollenberg, C. *et al.*, EP-A-0096430, GIST-BROCADES N.V., 1983).

The plasmids pUR2427 and pUR2428 are pTZ19R derivatives with the promoter and the DNA sequence encoding either the signal peptide (=pre-sequence) (in pUR2428), or the natural prepro-sequence (in pUR2427), of inulinase (inu) from Kluyveromyces marxianus. Both plasmids contain a unique BspMl site suitable to create a perfect joint with Eagl or Notl digested DNA-fragments (not yet published European patent application 92203932.6, supra). In both plasmids a unique HindIII site is located a bit further downstream of the BspMI-site, so that Eagl-HindIII cut DNA-fragments encoding V_{II} from Camelidae either solely or with Myc- or Flag- tail

can be easily ligated into BspM1-HindIII digested pUR2427 or pUR2428. Thereby a set of six plasmids can be created containing the promoter and secretion signals of the Kluyveromyces marxianus inulinase gene, joint in frame to Camelidae Vh encoding sequences, all on a EcoRI-HindIII restriction fragment:

The EcoRI-HindIII fragments of these plasmids can be ligated into the expression

vector pSK1 (not yet published European patent application 92203932.6, supra),

from which the α-galactosidase expression cassette including the GAL7-promoter is removed with a EcoRI(partial) and HindIII digestion. The resulting plasmids can then be transformed for example in K. lactis strain MSK110 (a, uraA, trp1::URA3), as they contain the trp1 marker and the pKD1 episomal plasmid sequences:

pUR4447 P_{inu} - Inu prepro seq. - V_H - 03 20 pUR4447M P_{inu} - Inu prepro seq. - V_H - 03 - Myc pUR4447F P_{inu} - Inu prepro seq. - V_H - 03 - Flag pUR4448 P_{inu} - Inu pre seq. - V_H - 03 - Myc pUR4448M P_{inu} - Inu pre seq. - V_H - 03 - Myc pUR4448F P_{inu} - Inu pre seq. - V_H - 03 - Flag .

25 A map of pUR4447 is shown in Figure 14.

Transformation can be performed by standard techniques such as the methods of Beggs (1978) or electroporation, using 0.67% Yeast Nitrogen Base (without amino acids) and 2% glucose as the selection medium for transformants.

Construction of Kluyveromyces lactis multicopy integration vectors. 4.2.

Alternatively, since all tailed and non-tailed versions of the Vh fragments, joined to the inulinase promoter and secretion signals, are located on EcoRI-HindIII fragments, the rDNA multicopy integration plasmid pMIRKGAL-Ta1 (Bergkamp et al., 1992) can be used in a similar way as the pSK1 plasmid. In order to replace the α-gal expression cassette present in this plasmid, by a antibody fragment cassette, these plasmids have to be digested with EcoRI(partial) and HindIII. After isolating the vector fragments, they can be ligated with the about 1.2 kb EcoRI-HindIII fragments which can be obtained from the plasmids described in example 4.1. The 10 resulting plasmids can be linearized with SacII and transformed to MSK110, resulting in K. lactis strains with potentially high and stable expression of single chain V₁₁ fragments.

pUR4449 P_{inu} - Inu prepro seq. - V_{II} - 03 pUR4449M P_{inu} - Inu prepro seq. - V₁₁ - 03 - Myc P_{inu} - Inu prepro seq. - V_H - 03 - Flag pUR4449F pUR4450 P_{inu} - Inu pre seq. - V_H - 03 pUR4450M P_{inu} - Inu pre seq. - V_H - 03 - Myc pUR4450F P_{inu} - Inu pre seq. - V_H - 03 - Flag.

20 4.3. Construction of Kluyveromyces marxianus episomal plasmids.

Kluyveromyces marxianus is a yeast which is perhaps even more attractive than K. lactis for industrial biotechnology, due to its short generation time on glucose (about 45 minutes) and its ability to grow on a wide range of substrates, and its growth at elevated temperatures (Rouwenhorst et al., 1988).

The shuttle vector pUR2434, containing the leu2 marker and the pKD1 plasmid sequences (not yet published European patent application 92203932.6, supra), located on a pUC19 based vector, can be cut with EcoRI(partial) and HindIII to remove the α-galactosidase expression cassette. In this vector the EcoRI-HindIII fragments containing the Vh expression cassettes as described in example 4.1, can be ligated. The resulting plasmids can then be transformed into KMS3, the neat leu2auxotroph CBS6556 K. marxianus strain (Bergkamp, 1993) using the method of Meilhoc et al. (1990).

```
pUR4451 P_{inu} - Inu prepro seq. - V_{II} - 03

pUR4451M P_{inu} - Inu prepro seq. - V_{II} - 03 - Myc

pUR4451F P_{inu} - Inu prepro seq. - V_{II} - 03 - Flag

pUR4452 P_{inu} - Inu pre seq. - V_{II} - 03 - Myc

pUR4452M P_{inu} - Inu pre seq. - V_{II} - 03 - Myc

pUR4452F P_{inu} - Inu pre seq. - V_{II} - 03 - Flag .

A map of pUR4451 is shown in Figure 15.
```

4.4 Construction of Kluyveromyces marxianus multicopy integration vectors.

- For high and stable expression in Kluyveromyces marxianus, the multicopy integration system as described by Bergkamp (1993), can be used. The following cloning route, based on the route for constructing pMIRKM-GAL5 (Bergkamp, 1993), results in suitable expression vectors for production of Vh fragments from Camelidae. The EcoRI-NheI(Klenow filled) fragments of pUR4447,-M,-F and pUR4448,-M,-F containing the Vh fragment expression cassettes as described in example 4.1, can be isolated and ligated in EcoRI-EcoRV digested pIC-20H. From the plasmids obtained in this way, and which are equivalents of the pIC-agal plasmid, the BamHI-NnII fragment can be isolated and ligated with BamHI-SmaI digested pMIRKM4. The result of this will be expression vectors which are equivalent to pMIRKM-GAL5, and contain a tailed or non-tailed Vh fragment from camel under control of inulinase promoter and secretion signals, in a vector which also contains the K marxianus LEU2-gene with defective promoter, and K. marxianus rDNA sequences for targeted integration into the genome. These vectors can be used to transform for example KMS3.

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Example 5. Construction of Hansenula polymorpha integrating vectors for the expression of (functionalized) V₁₁ fragments from *Camelidae*.

In search for productive systems able to carry out authentic posttranscriptional processing and overcoming the limitation of higher eukaryotic expression systems,

5 such as high costs, low productivity and the need for stringent control procedures for the detection of contaminating agents could be overcome by the methylotrophic yeast *H. polymorpha*. This strain is able to grow on methanol as its sole carbon and energy source, so the presence of methanol in the growth medium rapidly induces the enzymes of the methanol pathway, such as the key enzymes methanol oxidase

10 (MOX) and dihydroxyacetone synthase (DHAS).

While experiments to express foreign genetic information from an episomal plasmid resulted a low plasmid stability, chromosomal integration is the method of choice (Sierkstra et al., 1991). By utilizing the DNA of the mox gene as integration locus the latter were able to express and secrete α -galactosidase regulated by mox promoter and -terminator. Here, the S. cerevisiae SUC2 signal sequence was proven to be efficiently functional for secretion.

The same approach can be used for expression and secretion of Camelidae V_H antibody fragments. Plasmids analogous to pUR3515 (without an origin of replication functional in yeast) and pUR3517 (containing the HARS2 sequence as 20 origin of replication) can be used as expression vectors (Sierkstra et al., 1991). As a starting vector pUR3501 can be used (Sierkstra et al., 1991) in which by means of site directed mutagenesis (e.g. via PCR technology), an EagI restriction site is introduced at the junction between the invertase (=SUC2) signal sequence and the α-galactosidase. From the resulting plasmid, pUR3501Eag, it is possible to replace the Eagl-HindlII fragment comprising the α-galactosidase gene by an Eagl-HindIII fragment encoding a (functionalized) antibody fragment, obtained as described in example 1. In case of using the Eagl-HindIII fragments of the pUR4421-03 series (example 1), this would result in plasmids pUR4437 (Figure 17), pUR4437M and pUR4437F. In these plasmids the nucleotide sequence encoding the (functionalized) 30 V_{II} is preceded by a nucleotide sequence encoding the invertase signal sequence and the mox promoter sequence. The obtained plasmids can be digested with BamHI and HindIII and after filling in the sticky ends with Klenow polymerase, the about

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2.6 kb fragments can be ligated into plasmid pUR3511 which was digested with Smal (Sierkstra et al., 1991). In this way the terminator sequence of the mox gene can by fused downstream of the V_{II} encoding sequences. From the thus obtained plasmids, pUR4438 (Figure 17) EcoRI-HindIII fragments of about 3 kb can be isolated, containing the mox promoter, the invertase signal sequence, the (functionalized) V_{II} fragment and the mox transcription terminator. Subsequently these fragments can be cloned into plasmid pUR3513 (no yeast origin of replication) or in pUR3514 (HARS origin of replication) as described by Sierkstra et al. (1991), resulting in two sets of plasmids:

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```
pUR4439 P<sub>mox</sub> - SUC2 sig. seq. - V<sub>H</sub> - mox term. -- no origin
pUR4439M P<sub>mox</sub> - SUC2 sig. seq. - V<sub>H</sub> - mox term. -- no origin
pUR4439F P<sub>mox</sub> - SUC2 sig. seq. - V<sub>H</sub> - mox term. -- no origin
pUR4440 P<sub>mox</sub> - SUC2 sig. seq. - V<sub>H</sub> - mox term. -- HARS origin
pUR4440M P<sub>mox</sub> - SUC2 sig. seq. - V<sub>H</sub> - mox term. -- HARS origin
pUR4440F P<sub>mox</sub> - SUC2 sig. seq. - V<sub>H</sub> - mox term. -- HARS origin
Maps of pUR4439 and pUR4440 are shown in Figure 18.
```

Essentially the same can be done with other *EagI-HindIII* fragment, obtained as described in example 1.

The newly obtained plasmids can be transformed by electroporation of *H.* polymorpha A16 (CBS4732, leu-) and can be selected by growing on selective medium containing 0.68% YNB and 2% glucose. Induction medium should contain 0.5% methanol instead of the glucose.

25

Example 6 Construction Aspergillus niger var. awamori integration vectors for the production of V_{II} fragments from Camelidae.

The multiple cloning site of plasmid pEMBL9 (ranging from the *EcoRI* to the *HindIII* site) was replaced by a synthetic DNA fragment having the nucleotide sequence as indicated in Figure 19; see SEQ. ID. NO: 42-45. The 5'- part of the nucleotide sequence contains a *Nrul* restriction site followed by the first codons of the *Camelidae* V_{II} gene fragment and a *Xhol* restriction site. The 3'-part encodes for

a BstEII restriction site, the last codons of the Camelidae $V_{\rm H}$ gene, eleven codons of the Myc tail and finally a EcoRI and a AfIII site. The resulting plasmid is pUR4432.

After digesting plasmid pB3 with Xhol and EcoRI, a DNA fragment of approximately 425 bp can be isolated from agarose gel. This fragment codes for a truncated V_{II}-Flag fragment, missing the first 5 amino acids of the Camelidae V_{II}. The obtained fragment can be cloned into pUR4432. To this end plasmid pUR4432 can be digested with Xhol and EcoRI, after which the about 4 kb vector fragment was isolated from an agarose gel. Ligation with the about 425 bp fragment resulted in plasmid pUR4433F.

After digesting the plamids pB3 with XhoI and BstEII, a DNA fragment of approximately 365 bp was isolated from agarose gel. This fragment codes for a truncated $V_{\rm II}$ fragments, missing the first and last 5 amino acids of the Camelidae $V_{\rm H}$.

15 The obtained fragment was cloned into pUR4432. To this end plasmids pUR4432 can be digested with Xhol and BstEll, after which the about 4 kb vector fragment was isolated from an agarose gel. Ligation with the about 365 bp fragments resulted in plasmids pUR4433M. In a similar way the Xhol-BstEll fragments of pB9 and pB24 were cloned into the pUR4432 vector fragment, resulting in pUR4434M and pUR4435M, respectively.

Upon digesting pUR4433M or pUR4433F with BstEII and HindIII, the vector fragments of about 4.4 kb can be isolated from agarose gel and religated in the presence of a synthetic linker peptide having the following sequence:

25 Af III HindIII
25 GTCACCGTCTCCTCATAATGATCTTAAGGTGATA

GCAGAGGAGTATTACTAGAATTCCACTATTCGA (see SEQ. ID. NO: 50-51).

In the thus obtained plasmid, pUR4433, the Myc tail or Flag tail sequences are removed and the V_{II} gene fragment is directly followed by a stop codon.

Analogous as described in example 1.5, it is possible to clone nucleotide sequences encoding longer fragments of the heavy chain immunoglobulins into pUR4432 or to replace the BstEll-AfIII fragments of the above mentioned plasmids pUR4433,

pUR4433F or pUR4433M with other BstEll-AffII fragments, resulting in frame fusions encoding functionalized V_{II} fragments, having a C-terminal extension. Upon replacing the Nrul-XhoI fragments of pUR4433, pUR4433F or pUR4433M, in frame fusions can be constructed encoding functionalized V_{II} fragments, having an N-terminal extension.

In the above described constructs an Nrul site was introduced before the first codon of the (functionalized) V_{II} fragment, facilitating an in frame fusion with the precursor-sequence of xylanase, see (not prior-published) WO-A-93/12237, supra. For the construction of Aspergillus expression plasmids, from the plasmids pUR4433F, pUR4433M and pUR4433, respectively, an about 455, 445 and 405 bp Nrul-AfIII fragment has to be isolated encoding the V_{II} fragment with a Flag, a Myc or no tail.

Plasmid pAW14B was the starting vector for construction of a series of expression plasmids containing the exlA expression signals and the genes coding for (functionalized) V_H fragments of Camelidae heavy chain antibodies. The plasmid comprises an Aspergillus niger var. awamori chromosomal 5 kb SalI fragment on which the 0.7 kb exlA gene is located, together with 2.5 kb of 5'-flanking sequences and 2.0 kb of 3'-flanking sequences (see Figure 20 and (not prior-published) WO-A-20 93/12237, supra).

Starting from pAW14B, pAW14B-10 was constructed by removing the *Eco*RI site originating from the pUC19 polylinker, and introducing a *Not*I site. This was achieved by digesting plasmid pAW14B with *Eco*RI and after dephosphorylation the linear 7.9 kb *Eco*RI fragment was isolated. The fragment was religated in the presence of the "*Eco*RI"-*Not*I linker:

5'- AATTGCGGCCGC -3'

(see SEQ. ID. NO: 52).

Subsequently the AfIII site, located downstream of the exlA terminator was removed by partially cleaving plasmid pAW14B-10 and religating the isolated, linearized plasmid after filling in the sticky ends, resulting in plasmid

30 pAW14B-11.

25

Finally, pAW14B-12 was constructed using pAW14B-11 as starting material. After digestion of pAW14B-11 with AfIII (overlapping with the exlA stop codon) and BgIII

(located in the ext promoter) the ~2.4 kb AfIII-BgIII fragment, containing part of the extA promoter and the extA gene was isolated as well as the ~5.5 kb AfIII-BgIII vector fragment. After partial digestion of this ~2.4 kb fragment with BspHI (located in the extA promoter and at the extA start codon) an about 1.8 kb BgIII-BspHI extA promoter fragment (up to the ATG initiation codon) was isolated and ligated with the about 5.5 kb AfIII-Bg/II vector fragment of pAW14B-11 in the presence of the following adaptor:

(BspHI) BbsI AflII
CATGCAGTCTTCGGGC
GTCAGAAGCCCGAATT

(see SEQ. ID. NO: 53-54).

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30

For the construction of the V_{II} expression plasmids, pAW14B-11 can be partially digested with NruI and digested with AfIII, after which the ⁻ 7 kb vector fragment can be isolated from agarose gel and contains the xylanase promoter, the DNA sequence encoding the xylanase signal sequence and the xylanase terminator. Upon ligation of the NruI-AfIII fragments of pUR4433M, pUR4434M and pUR4435M with the pAW14B-11 vector, plasmids pUR4436M, pUR4437M and pUR4438M were obtained, respectively. In these plasmids the Camelidae V_H polypeptides are preceded by the 27 amino acid long precursor sequence of xylanase and followed by the myc-tail (of 11 amino acids; see Examples 1.3 en 2, Figures 6 and 19, and

In a similar way plasmids can be constructed encoding the V_H fragments followed by the FLAG-tail or without a tail.

After introducing the amdS and pyrG selection markers into the unique NotI site of pUR4436M, pUR4437M and pUR4438M using conventional techniques, e.g. as described in Examples 2 and 3 of (not prior-published) WO-A-93/12237, supra, the

plasmids were transferred to Aspergillus.

SEQ.ID. NO: 41 = 45).

Production of the Camel V₁₁ fragments by the selected transformants was achieved by growing the strains in inducing medium essentially as described in example 2.2 of (not prior-published) WO-A-93/12237, *supra*. Western blot analysis of the culture medium was perforemed as described in Example 2.1 above and revealed the presence of the antibody fragments.

Obviously, expression vectors can be constructed in which different promoter systems, e.g. glucoamylase promoter, and/or different signal sequences, e.g. glucoamylase or glucose oxidase signal sequences, are used.

- 5 Example 7 Production of glucose oxidase V_{II} fusion proteins
 - Glucose oxidase catalyses the oxidation of D-glucose to D-gluconate under the release of hydrogen peroxide. Glucose oxidase genes (gox) from Aspergillus niger have been cloned (Frederick et al. (1990) J. Biol. Chem. 265 3793, Kriechbaum et al., 1989) and the nucleotide sequences are available from the EMBL data bank
- 10 under accession numbers J05242 and X16061. The nucleotide sequence of the latter is used as a basis for the following construction route.
 - Upon cloning the gox gene from A. niger it is possible, by applying PCR technology, to introduce convenient restriction sites.
 - To introduce a BspHI restriction site, overlapping with the ATG initiation codon, the sequence ATC ATG CAG can be changed to ATC ATG AGG. In the same experiment an EcoRI restriction site can be introduced which is located upstream of the BspHI site. This can be achieved by using the following PCR primer:
 - EcoRI BspHI
 5'-TCACTGAATTCGGGATC ATG AGG ACT CTC CTT GTG AGC TCG CTT-3'
 (see SEQ. ID. NO: 55).

A second PCR primer, having the following sequence can be used:

- AflI BbsI SalI
 5'-ATGTCACAAAGCTTAAGCACGAAGACA GTC GAC CGT GCG GCC GGA GAC-3'
 HindII
- 25 (see SEQ. ID. NO: 56)
 - in the same PCR experiment, in order to introduce a BbsI site, a AfIII site and a HindIII site, downstream of the unique SalI site present in the glucose oxidase gene. After digesting the DNA obtained from this PCR experiment with EcoRI and HindIII, an EcoRI HindIII fragment of about 160 bp can be isolated and cloned
- into pEMBL9, which was digested with the same enzymes, resulting in plasmid pGOX1.
 - From pGOX1 an about 140 bp BspHI AfIII fragment can be isolated and introduced into the 7.2 kb BbsI-AfIII vector fragment of pAW14B-12, resulting in

25

pAW14B-GOX. In this plasmid, the 5'- part of the gox gene, encoding the first 43 amino acids, is fused in frame with the ATG initiation codon of the exlA gene.

In a second PCR experiment, a Mlul restriction site can be introduced near the 3'end of the gox by changing the sequence TAT GCT TCC to TAC GCG TCC. In the same experiment a HindlII site can be introduced downstream of the MluI site. As a second primer an oligo nucleotide should be used hybridizing upstream of the Sall site. After digesting the DNA obtained from this PCR experiment with Sall and HindIII, an Sall - HindIII fragment of about 1.7 kb can be isolated and cloned into 10 pEMBL9, which was digested with the same enzymes, resulting in plasmid pGOX2. Upon digesting pGOX2 with MluI and HindIII, an about 5.7 kb vector fragment can be isolated.

From the plasmids pUR4433, pUR4433F, pUR4433M and the like, XhoI-HindIII fragments can be isolated, encoding the truncated Camelidae V_{II} fragment with or 15 without a tail sequence, and missing the first 4-6 N-terminal amino acids (see Example 1). These fragments can be ligated into the 5.7 kb pGOX2 vector fragment by using MluI-XhoI adaptors. These adaptors are designed in such a way that they result in an in frame fusion between the 3'-end of the gox gene and the restored V_H gene fragment, optionally intersected with a DNA sequence encoding a peptide linker sequence.

An example of these designed adaptors is:

MluI XhoT CGCGTCCATGCAGTCCTCAGGTGGATCATCCCAGGTGAAACTGC AGGTACGTCAGGAGTCCACCTAGTAGGGTCCACTTTGACGAGCT S G G S S Q V K L L E S M Q S (see SEQ. ID. NO: 57-59)

which encodes for the last amino acids of GOX, an SSGGSS linker sequence (see ... SEQ. ID. NO: 62) and the N-terminal amino acids of the Camel V_H fragment of pB3. Instead of the SSGGSS linker (see SEQ. ID. NO: 62) it is possible to use other linkers such as the repeated sequence linkers described in the above indicated European patent application 92402326.0, e.g. a repeated sequence Pro-X, with X being any amino acid, but preferably Gln, Lys or Glu, the sequence containing

advantageously at least 3 repeats of Pro-X and especially a fragment composed of a 12-fold repeat of the sequence Pro-X.

In case the about 435 bp Xhol-HindIII fragment of pUR4433M is used in combination with the above described adaptor, this would result in pGOX2-03M. From this plasmid a Sall-AfIII fragment of about 2.1 kb encoding the C-terminal part of glucose oxidase followed by the linker peptide, the Camel V_{II} fragment of pB3 and finally the Myc tail.

Upon digesting pAW14B-GOX partially with BbsI, and with AfIII, the about 7.4 kb vector fragment can be isolated. This fragment contains the xylanase promoter, the DNA sequence encoding the N-terminal part of glucose oxidase and the xylanase promoter. Due to the digestion with BbsI, a SalI sticky end is created, corresponding with the SalI restriction site originally present in the gox gene. Ligation of the SalI-AfIII vector fragment with the about 2.1 kb SalI-AfIII fragment of pGOX2-03M,

resulting in pUR4441M. This expression plasmid encodes for a single chain polypeptide comprising the glucose oxidase enzyme, the (functionalized) Camel V_H fragment and the Myc tail.

Introduction of this type of expression plasmids in Aspergillus can be achieved essentially as described in example 6.

As the naturally occurring glucose oxidase is a homodimeric enzyme, it might be expected that a fusion protein, comprising glucose oxidase and an antibody fragment as a C-terminal extension, has an increased avidity for the antigen/antibody binding, if this fusion protein is produced as a homodimer. Alternatively, it is possible to produce heterodimers, consisting of one glucose oxidase molecule connected to a V_H fragment and one wild type glucose oxidase molecule. This can be achieved by producing with the same strain both wild type glucose oxidase and the fused glucose oxidase-V_H fragment, or by mixing the two different homodimers produced by different strains under conditions whereby the mixture of dimers are dissociated and subsequently associated.

Example 8 Engineering of Camelidae V₁₁ fragments

8.1 Random and targeted random mutagenesis.

After expressing a number of different Camelidae V_H fragments in lower eukaryotic host organisms as described above, or in prokaryotes, fragments produced in relative higher amounts can be selected. Upon subjecting the Xhol-BstEII gene fragments to a (targeted) random mutagenesis procedure, it might be possible to further improve special characteristics of the V_H fragment, e.g. further improvement of the production level, increased stability or increased affinity.

To this end the following procedure might be followed.

described in previous examples.

10 Upon replacing the polylinker of the phagemid vector pHEN1 (Hoogenboom et al., 1991) located on a Ncol-Notl fragment by a new polylinker having the following sequence:

NCOI XhOI BSTEII NOTI
CATGGCCAGGTGAAACTGCTCGAGTAAGTGACTAAGGTCACCGTCTCCTCAGC
CGGTCCACTTTGACGAGCTCATTCACTGATTCCAGTGGCAGAGGAGTCGCCGG

(see SEQ. ID. NO: 60-61) it becomes possible to introduce XhoI-BstEII fragments encoding truncated Camelidae V_H fragments in the phagemid.

Following mutagenesis of the V_H encoding sequence (random mutagenesis) or a specific part thereof (targeted random mutagenesis), the mutated V_H fragments can be expressed and displayed on the phage surface in essentially the same way as described by Hoogenboom et al. (1991). Selecting phages displaying (mutant) V_H fragments, can be done in different ways, a number of which are described by Marks et al. (1992). Subsequently, the mutated XhoI-BstEII fragments can be isolated from the phagemid and introduced into expression plasmids for yeast or fungi as

Upon producing the mutant V_{II} fragments by these organisms, the effects of the mutations on production levels, V_{II} fragment stability or binding affinity can be evaluated easily and improved V_{II} fragments can be selected.

Obviously, a similar route can be followed for larger antibody fragments. With similar procedures the activity of catalytic antibodies can be improved.

8.2 Site-directed or designed mutagenesis

As an alternative to the methods described above in Example 8.1 it is possible to use the well-known technique of site-directed mutagenesis. Thus, designed mutations, preferably based on molecular modelling and molecular dynamics, can be introduced in the V₁₁ fragments, e.g. in the framework or in the CDRs.

8.3 Construction V_{II} fragments with regulatable binding efficiencies.

For particular applications, the possibility to regulate the binding capacity of antibody fragments might be necessary. The introduction of metal ion binding sites in proteins is known from the literature e.g. Pessi et al. (1993). The present inventors envisage that the introduction of a metal binding site in an antibody fragment by rational design can result in a regulatable antibody fragment, when the metal binding site is introduced at a position such that the actual binding of the metal ion results in a conformational change in the antibody fragments due to which the binding of the antigen to the antibody fragment is influenced. Another possibility is that the presence of the metal prevents antigen binding due to steric hindrance.

8.4 Grafting of CDR regions on the framework fragments of a Camelidae V_H fragment.

- Grafting of CDR fragments onto framework fragments of different antibodies or fragments thereof is known from the literature (see Jones et al. (1986), WO-A-92/15683, and WO-A-92/01059). In these cases the CDR fragments of murine antibody fragments were grafted onto framework fragments of human antibodies. The sole rationale behind the "humanization" was to increase the acceptability for therapeutic and/or diagnostic applications in human.
 - Essentially the same approach can however also be used for a totally different purpose. Although antibody fragments share some homology in the framework areas, the production levels vary considerably.
- Once an antibody or an antibody fragment, e.g. a *Camelidae* V_{II} fragment, has been identified, which can be produced to high levels by an production organism of interest, this antibody (fragment) can be used as a starting point to construct "grafted" antibody (fragments), which can be produced in high levels and have an

other specificity as compared to the original antibody (fragment). In particular cases it might be necessary to introduce some modifications in the framework fragments as well in order to obtain optimal transitions between the framework fragments and the CDR fragments. For the determination of the optimal transitions molecular dynamics and molecular modelling can be used.

To this end a synthetic gene, encoding the "grafted V_{11} " fragment, can be constructed and introduced into an expression plasmid. Obviously it is possible to adapt the codon usage to the codons preferred by the host organism.

For optimization of the "grafted V_{II} " fragment, the procedure as described in example 8.1 can be followed.

Literature mentioned in the specification additional to that mentioned in the above given draft publication

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 - Harmsen, M.M., Langedijk, A.C., van Tuinen, E., Geerse, R.H., Rauè, H.A., Maat, J., (1993) Effect of pmr1 disruption and different signal sequences on the

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 - Sierkstra, L.N., Verbakel, J.M.A. and Verrips, C.T. (1991) Optimisation of a host/vector system for heterologous gene expression by *Hansenula polymorpha*.
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 - Teeri et al., WO-A-93/02198 (TECH. RES. CENT. FINLAND, publ. 04.02.1993)
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Additional references to prior-filed but not prior-published patent applications, which are incorporated herein by reference:

- not prior-published PCT application EP 92/02896, filed 09.12.92 with priority date of 09.12.91 (UNILEVER / R.J. Gouka et al.), now publicly available as WO-A-93/12237
- not prior-published EP application 92202080.5, filed <u>08.07.92</u> (UNILEVER / F.M. Klis et al.), now publicly available as International (PCT) patent application WO-A-94/01567)
- not prior-published EP application 92402326.0, filed <u>21.08.92</u> (C. Casterman & R.
 Hamers), now publicly available as EP-A1-0 584 421
 - not yet published EP application 92203932.6, filed 11.12.92 (UNILEVER / H.Y. Toschka & J.M.A. Verbakel).

15

5

Information on deposits of micro-organisms under the Budapest Treaty is given in Example 1 on page 23, lines 23-25 above. In agreement with Rule 28 (4) EPC, or a similar arrangement for a State not being a Contracting State of the EPC, it is hereby requested that a sample of such deposit, when requested, will be submitted to an expert only.

SEQUENCE LISTING

```
(1) GENERAL INFORMATION:
 5
           (i) APPLICANT:
                 (A) NAME: Unilever N.V.
                 (B) STREET: Weena 455 (C) CITY: Rotterdam
                 (E) COUNTRY: The Netherlands
10
                 (F) POSTAL CODE (ZIP): NL-3013 AL
                 (A) NAME: Unilever PLC
                 (B) STREET: Unilever House Blackfriars
                 (C) CITY: London
15
                 (E) COUNTRY: United Kingdom (F) POSTAL CODE (ZIP): EC4P 4BQ
                 (A) NAME: Leon Gerardus Joseph FRENKEN
                 (B) STREET: Geldersestraat 90
20
                 (C) CITY: Rotterdam
                 (E) COUNTRY: The Netherlands
                 (F) POSTAL CODE (ZIP): NL-3011 MP
                 (A) NAME: Cornelis Theodorus VERRIPS
25
                 (B) STREET: Hagedoorn 18
                 (C) CITY: Maassluis
                 (E) COUNTRY: The Netherlands
                 (F) POSTAL CODE (ZIP): NL-3142 KB
30
                 (A) NAME: Raymond HAMERS
                 (B) STREET: Vijversweg 15
(C) CITY: Sint-Genesius-Rode
                 (E) COUNTRY: Belgium
                 (F) POSTAL CODE (ZIP): B-1640
35
                 (A) NAME: Cécile HAMERS-CASTERMAN
                (B) STREET: Vijversweg 15
                (C) CITY: Sint-Genesius-Rode
                 (E) COUNTRY: Belgium
40
                 (F) POSTAL CODE (ZIP): B-1640
                (A) NAME: Serge Victor Marie MUYLDERMANS
                (B) STREET: Brusselse Steenweg 55 (C) CITY: Hoeilaart
45
                 (E) COUNTRY: Belgium
                 (F) POSTAL CODE (ZIP): B-1560
          (ii) TITLE OF INVENTION: Production of antibodies or (functionalized)
                   fragments thereof derived from heavy chain immunoglobulins
50
                  of Camelidae.
        (iii) NUMBER OF SEQUENCES: 62
          (iv) COMPUTER READABLE FORM:
55
                (A) MEDIUM TYPE: Floppy disk
                (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
                (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
60
     (2) INFORMATION FOR SEQ ID NO: 1:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 5 amino acids (B) TYPE: amino acid
65
                (C) STRANDEDNESS: single
```

(D) TOPOLOGY: linear

	(ii)	MOLECULE	TYPE:	protei	n	٠								
	(xi)	SEQUENCE	DESCRI	PTION:	SEQ II	NO:	1:							
5	Ala: 1	Pro Glu I	Leu Leu 5											
10	(2) INFOR													
15	(1)	SEQUENCE (A) LENG (B) TYPE (C) STRA (D) TOPG	E: amino ANDEDNE:	amino a o acid SS: si	acids									
	(ii)	MOLECULE	TYPE: 1	protei	n									
20	(xi)	SEQUENCE	DESCRI	PTION:	SEQ II	NO:	2:							
20	Ala 1	Pro Glu I	Leu Pro 5											
25	(2) INFOR	MATION FO	OR SEQ	ID NO:	3:									
30	(1,)	SEQUENCE (A) LENG (B) TYPE (C) STR	STH: 21 E: nucle Andedne:	base peic ac: SS: sin	pairs id									ì
		(D) TOPO												
25	(ii)	MOLECULE	TYPE: 1	DNA (ge	enomic))								
35	(xi)	SEQUENCE	DESCRI	PTION:	SEQ II	NO:	3:		-					
	CGCCATCAA	G GTACCAC	STTG A											21
40	(2) INFOR	MATION FO	OR SEQ	ID NO:	4:									
45	(i)	SEQUENCE (A) LENG (B) TYPI (C) STRI (D) TOPG	GTH: 89 E: amin ANDEDNE:	amino o acid SS: si	acids									
5 0	(ii)	MOLECULE	TYPE:	protei	n									
50	(vii)	IMMEDIATI (B) CLO	NE: hum	an hea	vy cha: R1, Xaa								= CDF	t3) ·
55	(xi)	SEQUENCE	DESCRI	PTION:	SEQ II	NO:	4:							
	Glu 1	Val Gln 1	Leu Val 5	Glu S	er Gly	Gly	Gly 10	Leu	Val	Gln	Pro	Gly 15	Gly	
60	Ser	Leu Arg l	Leu Ser 20	Cys A	la Ala	Ser 25	Gly	Xaa	Trp	V al	Arg 30	Gln	Ala	
65	Pro	Gly Lys (Gly Leu	Glu T	rp Val 40	Ser	Xaa	Xaa	Arg	Phe 45	Thr	Ile	Ser	
V.J	Arg	Asp Asn S	Ser Lys	Asn T	hr Leu	Tyr	Leu	Gln	Met	Asn	Ser	Leu	Arg	

	Ala 65	Glu	Asp	Thr	Ala	Val 70	Tyr	Tyr	Cys	Ala	Arg 75	Xaa	Xaa	Xaa	Trp	.Gly 80
5	Gln	Gly	Thr	Leu	Val 85	Thr	Val	Ser	Ser							
	(2) INFO	RMAT	ION :	FOR :	SEQ :	ID N	0: 5	:								
10	(i)	(A (B (C) LEI) TYI) STI	NGTH PE: (RAND)	ARAC' : 81 amino EDNE:	ami o ac. SS: :	no a id sing	cids								
15	(ii)															
20	(vii)				came	el "1	heav CDR1	y ch	ain : a Xa	immu: a = (nogli CDR2	obul and	in" Xaa	fram Xaa	ewor: Xaa	k A = CDR3)
	(xi)	SEQ	UENCI	E DES	SCRII	PTIO	N: S	EQ I	D NO:	: 5:		٠				
25	Gly 1	Gly	Ser	Val	Gln 5	Gly	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Сув	Ala 15	Ile
	Ser	Gly	Xaa	Trp 20	Phe	Arg	Glu	Gly	Pro 25	Gly	Lys	Glu	Arg	Glu 30	Gly	Ile
30	Ala	Xaa	Xaa 35	Arg	Phe	Thr	Ile	Ser 40	Gln	Asp	Ser	Thr	Leu 45	Lys	Thr	Met
35	Tyr	Leu 50	Leu	Met	Asn	Asņ	Leu 55	Lys	Pro	Glu	Asp	Thr 60	Gly	Thr	Tyr	Tyr
	Сув 65	Ala	Ala	Xaa	Xaa	Xa a 70	Trp	Gly	Gln	Gly	Thr 75	Gln	Val	Thr	Val	Ser 80
40	Ser															
	(2) INFO	RMAT	ON I	FOR S	SEQ 1	D NO): 6:	:								
4 5	(i)	(A) (B) (C)	LEN TYP STF	NGTH: PE: & RANDI	ARACT 81 amino EDNES SY:]	amir o aci SS: s	no ad id sing!	cids								
50	(ii)	MOLE	CULE	TYI	PE: p	prote	ein									
	(Vii)				came	el "r	neavy	y cha . Xaa	ain i a Xaa	immur a = C	noglo	bul:	in" : Xaa	frame Xaa	ewor) Xaa	c B = CDR3)
55	(xi)	SEQU	JENCE	E DES	CRIE	OIT	1: SI	EQ II) NO:	6:						·
50	Gly 1	Gly	Ser	Val	Gln 5	Ala	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Cys	Ala 15	Ser
	Ser	Ser	Xaa	Trp 20	Tyr	Arg	Gln	Ala	Pro 25	Gly	Lys	Glu	Arg	Glu 30	Phe	Val
55	Ser	Xaa	Xaa 35	Arg	Phe	Thr	Ile	Ser 40	Gln	Asp	Ser	Ala	Lys 45	Asn	Thr	Val

	Tyr	Leu Gln M 50	let Asn	Ser Leu 55	Lys Pro	Glu Asp	Thr Ala	Met Tyr	Tyr
5	Сув 65	Lys Ile X	(aa Xaa	Xaa Trp 70	Gly Gln	Gly Thr 75	Gln Val	Thr Val	ser 80
	Ser								
10	(2) INFO	RMATION FO	OR SEQ I	D NO: 7:	:				
15	(i)	(B) TYPE	TH: 37 E: amino ANDEDNES	amino ad acid SS: singl	cids				
	(ii)	MOLECULE	TYPE: p	rotein			*		
20	(vii)	(B) CLON	ME: came	el "heavy			obulin" H2 fragm	ent	
25	(xi)	SEQUENCE	DESCRIE	PTION: SE	EQ ID NO	: 7:			
	Trp 1	Gly Gln G	Sly Thr 5	Gln Val	Thr Val	Ser Ser 10	Gly Thr	Asn Glu 15	Val
30	Cys	Lys Cys F	Pro Lys 20	Cys Pro	Ala Pro 25	Glu Leu	Pro Gly	Gly Pro 30	Ser
	Val	Phe Val I 35	Phe Pro						
35	(2) INFO	RMATION FO	OR SEQ I	ID NO: 8	:				
40	(1)	(B) TYPE	STH: 60 E: amino ANDEDNES	amino ac acid SS: sing	cids			·	
45	(ii)	MOLECULE	TYPE: I	protein					
	(vii)	(B) CLON	WE: came	el "heavy	y chain long hi	immunogl nge - CH	obulin" 2 fragme:	nt	
50	(xi)	SEQUENCE	DESCRI	PTION: S	EQ ID NO	: 8:			
	Trp 1	Gly Gln (Gly Thr 5	Gln Val	Thr Val	Ser Ser 10	Glu Pro	Lys Ile 15	Pro
55	Gln	Pro Gln I	Pro Lys 20	Pro Gln	Pro Gln 25	Pro Gln	Pro Gln	Pro Lys 30	Pro
60	Gln	Pro Lys 1 35	Pro Glu	Pro Glu	Cys Thr 40	Cys Pro	Lys Cys 45	Pro Ala	Pro
	Glu	Leu Leu (50	Gly Gly	Pro Ser 55	Val Phe	Ile Phe	Pro 60		

	(2) INFO	RMATION FOR SEQ ID NO: 9:	
5	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 67 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10		MOLECULE TYPE: protein	
	(A11)	IMMEDIATE SOURCE: (B) CLONE: human gamma-3 CH1 - hinge - CH2 fragment	
15		SEQUENCE DESCRIPTION: SEQ ID NO: 9:	
	Lys 1	Val Asp Lys Arg Val Glu Leu Lys Thr Pro Leu Gly Asp Thr Th 5 10 15	r
20	His	Thr Cys Pro Arg Cys Pro Glu Pro Lys Cys Ser Asp Thr Pro Pr 20 25 30	0
	Pro	Cys Pro Arg Cys Pro Glu Pro Lys Ser Cys Asp Thr Pro Pro Pro 35 40 45	0
2 5	Сув	Pro Arg Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Ph 50 60	e
30	Leu 65	Phe Pro	
	(2) INFO	RMATION FOR SEQ ID NO: 10:	
3 5	(1)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 35 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii)	MOLECULE TYPE: protein	
	(vii)	IMMEDIATE SOURCE: (B) CLONE: human gamma-1 CH1 - hinge - CH2 fragment	
45	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 10:	
	Lys 1	Val Asp Lys Lys Ala Glu Pro Lys Ser Cys Asp Lys Thr His Th	r
50	Сув	Pro Pro Cys Pro Ala Pro Glu Leu Leu Gly Gly Pro Ser Val Pho 20 25 30	е
55	Leu	Phe Pro 35	
	(2) INFO	RMATION FOR SEQ ID NO: 11:	
60	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 31 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
65	(ii)	MOLECULE TYPE: protein	

	(vii)	IMMEDIAT (B) CLO			-2 CH1 -	hinge -	СН2 1	fragment	
5	(xi)	SEQUENCE	DESCRI	PTION: S	EQ ID NO	: 11:			
•,	Lys 1	Val Lys	Val Thr 5	Val Glu	Arg Lys	Cys Cys 10	Val (Glu Cys	Pro Pro 15
10	Cys	Pro Ala	Pro Pro 20	Val Ala	Gly Pro 25	Ser Val	Phe I	Leu Phe 30	Pro
	(2) INFO	RMATION F	OR SEQ 1	ID NO: 1	2:				
15 20	(i)	(B) TYP (C) STR	GTH: 32 E: amino	amino a cacid SS: sing	cids				
2()	(ii)	MOLECULE	TYPE: p	protein					
25	(vii)	IMMEDIAT			-4 CH1 -	hinge -	CH2 1	Eragment	<u>:</u>
	(xi)	SEQUENCE	DESCRI	PTION: S	EQ ID NO	: 12:			
30	Lys 1	Val Asp	Lys Arg 5	Val Glu	Ser Lys	Tyr Gly 10	Pro I	Pro Cys	Pro Sei
	Cys	Pro Ala	Pro Glu 20	Phe Leu	Gly Gly 25	Pro Ser	Val I	Phe Leu 30	Phe Pro
35	(2) INFO	RMATION F	OR SEQ 1	ID NO: 1	3:	,			
40	(i)	(B) TYP (C) STR	GTH: 121 E: amino	l amino a cacid SS: sing	acids				
	(ii)	MOLECULE	TYPE: F	protein					
4 5	(vii)	IMMEDIAT		E: se heavy	chain V	-region			
	, ,	SEQUENCE							
50	1	Val Lys	5			10			15
55	Ser	Leu Arg	Leu Ser 20	Cys Ala	Thr Ser 25	Gly Phe	Thr I	Phe Ser 30	Asp Phe
	Tyr	Met Glu 35	Trp Val	Arg Gln	Pro Pro 40	Gly Lys		Leu Glu 45	Trp Ile
50	Ala	Ala Ser 50	Arg Asn	Lys Ala 55	Asn Asp	Tyr Thr	Thr 0	Glu Tyr	Ser Ala
	Ser 65	Val Lys	Gly Arg	Phe Ile 70	Val Ser	Arg Asp 75	Thr S	Ser Gln	Ser Ile 80
55	Leu	Tyr Leu	Gln Met 85	Asn Ala	Leu Arg	Ala Glu 90	Asp 3	Thr Ala	Ile Tyr 95

Tyr Cys Ala Arg Asp Tyr Tyr Gly Ser Ser Tyr Phe Asp Val Trp Gly Ala Gly Thr Thr Val Thr Val Ser Ser 5 (2) INFORMATION FOR SEQ ID NO: 14: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 131 amino acids (B) TYPE: amino acid(C) STRANDEDNESS: single (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: protein (vii) IMMEDIATE SOURCE: (B) CLONE: human heavy chain V-region 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14: Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly Gly 25 Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Phe Thr Phe Ser Ser Tyr Ala Met Ser Trp Val Arg Gln Ala Pro Gly Lys Gly Leu Glu Trp Val 30 Ser Xaa Ile Ser Xaa Lys Thr Asp Gly Gly Xaa Thr Tyr Tyr Ala Asp 35 Ser Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ser Lys Asn Thr 65 Leu Tyr Leu Gln Met Asn Ser Leu Arg Ala Glu Asp Thr Ala Val Tyr 40 Tyr Tyr Tyr His Xaa Phe Asp Tyr Trp Gly Gln Gly Thr Leu Val Thr 45 Val Ser Ser 130 50 (2) INFORMATION FOR SEQ ID NO: 15: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 114 amino acids 55 (B) TYPE: amino acid(C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein 60 -(vii) IMMEDIATE SOURCE: (B) CLONE: camel "heavy chain immunoglobulin" V-region (1) (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15: 65 Gly Gly Ser Val Gln Ala Gly Gly Ser Leu Arg Leu Ser Cys Ala Ala 10

						•										
	Ser	Gly	Tyr	Ser 20	Asn	Cys	Pro	Leu	Thr 25	Trp	Ser	Trp	Tyr	Arg 30	Glņ	Ph
5	Pro	Gly	Thr 35	Glu	Arg	Glu	Phe	Val 40	Ser	Ser	Met	Asp	Pro 45	Asp	Gly	As
	Thr	Lys 50	Tyr	Thr	Tyr	Ser	Val 55	Lys	Gly	Arg	Phe	Thr 60	Met	Ser	Arg	G1
10	Ser 65	Thr	Glu	Tyr	Thr	Val 70	Phe	Leu	Gln	Met	Asp 75	Asn	Leu	Lys	Pro	G1 80
15	Asp	Thr	Ala	Met	Tyr 85	Tyr	Cys	Lys	Thr	Ala 90	Leu	Gln	Pro	Gly	Gly 95	Ty
••	Cys	Gly	Tyr	Gly 100	Xaa	Cys	Leu	Trp	Gly 105	Gln	Gly	Thr	Gln	Val 110	Thr	Va
20	Ser	Ser														
	(2) INFO	RMAT:	I NO	FOR S	SEQ :	ID NO): 1 (5:								
25	(i)	(A) (B) (C)) LEI) TYI) STI	E CHI NGTH: PE: & RANDI POLOG	: 120 amino EDNES	ami aci	ino a id sing:	cids	3					-	,	
30	(ii)	MOLI	ECULI	E TYI	PE: 1	prote	ein									
	(vii)					_	neavy	y cha	ain i	i m mu r	oglo	buli	in" T	/-red	gion	(2)
35	(xi)	SEQ	JENCI	E DES	CRII	PTIO	1: SI	EQ II	NO:	16:	:					
	Asp 1	Val	Gln	Leu	Val 5	Ala	Ser	Gly	Gly	Gly 10	Ser	Val	Gln	Ala	Gly 15	Gl ₃
10	Ser	Leu	Arg	Leu 20	Ser	Cys	Thr	Ala	Ser 25	Gly	Asp	Ser	Phe	ser 30	Arg	Phe
15	Ala	Met.	ser 35	Trp	Phe	Arg	Gln	Ala 40	Pro	Gly	Lys	Glu	Суя 45	Glu	Leu	Va.
	Ser	Ser 50	Ile	Gln	Ser	Asn	Gly 55	Arg	Thr	Thr	Glu	Ala 60	Авр	Ser	Val	Glı
50	65					70		_			75				Tyr	80
	Gln	Met	Asn	Ser	Leu 85	Lys	Pro	Glu	Asp	Thr 90	Ala	Val	Tyr	Tyr	Cys 95	Gly
55	Ala	Val	Ser	Leu 100	Met	Asp	Arg	Ile	Ser 105	Gln	His	Gly	Cys	Arg 110	Gly	Glı
	Gly	Thr	Gln 115	Val	Thr	Val	Ser	Leu 120								
60																
50	(2) INFO	RMATI	ON I	FOR S	SEQ I	D NC): 17	7:								

		(D) TO	POLO	GY: :	line	ar									
	(ii)	MOL	ECULI	E TYI	PE:	prot	ein									
5	(vii)			TE SO			heav	y cha	ain :	immu	noglo	obul.	in" '	V-re	gion	(3)
	(xi)	SEQ	UENCI	E DES	SCRI	PTIO	N: SI	EQ II	NO.	: 17:	:					
10	Gly 1	Gly	Ser	Val	Gln 5	Thr	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Cys	Ala 15	Val
15	Ser	Gly	Phe	Ser 20	Phe	Ser	Thr	Ser	Cys 25	Met	Ala	Trp	Phe	Arg 30	Gln	Ala
••	Ser	Gly	Lys 35	Gln	Arg	Glu	Gly	Val 40	Ala	Ala	Ile	Asn	Ser 45	Gly	Gly	Gly
20	Arg	Thr 50	Tyr	Tyr	Asn	Thr	Tyr 55	Val	Ala	Glu	Ser	Val 60	Lys	Gly	Arg	Phe
	Ala 65	Ile	Ser	Gln	Asp	Asn 70	Ala	Lys	Thr	Thr	Val 75	Tyr	Leu	Asp	Met	Asn 80
25	•			Pro	85					90					95	
30	Ala	His	Leu	Gly 100	Pro	Gly	Ala	Ile	Leu 105	Asp	Leu	Lys	Lys	Tyr 110	Lys	Tyr
	Trp	Gly	Gln 115	Gly	Thr	Gln	Val	Thr 120	Val	Ser	Ser					
35	(2) INFO	RMAT:	ION 1	FOR S	SEQ I	ID NO): 18	3:								
40	(i)	(A) (B) (C)	LEI TYI STI	E CHA NGTH: PE: & RANDI POLOC	: 116 amino EDNES	sam: Sac:	ino a id sing]	acida	3.							
	(ii)	MOLI	ECULI	E TYI	PE: p	prote	ein									
45	(Vii)			TE SO ONE:			neavy	, cha	ain i	Lmmur	oglo	buli	in" 1	/-re	gion	(7)
	(xi)	SEQ	JENCI	E DES	SCRII	PTIO	N: SI	EQ II	ONO:	: 18:	;					
50	Gly 1	Gly	Ser	Val	Gln 5	Gly	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Сув	Ala 15	Ile
55	Ser	Gly	Tyr	Thr 20	Tyr	Gly	Ser	Phe	Cys 25	Met	Gly	Trp	Phe	Arg 30	Glu	Gly
	Pro	Gly	Lys 35	Glu	Arg	Glu	Gly	Ile 40	Ala	Thr	Ile	Leu	Asn 45	Gly	Gly	Thr
60	Asn	Thr 50	Tyr	Tyr	Ala	Asp	Ser 55	Val	Lys	Gly	Arg	Phe 60	Thr	Ile	Ser	Gln
	Asp 65	Ser	Thr	Leu	Lys	Thr 70	Met	Tyr	Leu	Leu	Met 75	Asn	Asn	Leu	Lys	Pro 80
65	Glu	Asp	Thr	Gly	Thr 85	Tyr	Tyr	Cys	Ala	Ala 90	Glu	Leu	Ser	Gly	Gly 95	Ser

55

Cys Glu L u Pro Leu Leu Phe Asp Tyr Trp Gly Gln Gly Thr Gln Val Thr Val Ser Ser 5 115 (2) INFORMATION FOR SEQ ID NO: 19: 10 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 114 amino acids (B) TYPE: amino acid
(C) STRANDEDNESS: single (D) TOPOLOGY: linear 15 (ii) MOLECULE TYPE: protein (vii) IMMEDIATE SOURCE: (B) CLONE: camel "heavy chain immunoglobulin" V-region (9) 20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19: Gly Gly Ser Val Gln Ala Gly Gly Ser Leu Thr Leu Ser Cys Val Tyr 25 Thr Asn Asp Thr Gly Thr Met Gly Trp Phe Arg Gln Ala Pro Gly Lys 20 25 30 Glu Cys Glu Arg Val Ala His Ile Thr Pro Asp Gly Met Thr Phe Ile 30 Asp Glu Pro Val Lys Gly Arg Phe Thr Ile Ser Arg Asp Asn Ala Gln 35 Lys Thr Leu Ser Leu Arg Met Asn Ser Leu Arg Pro Glu Asp Thr Ala Val Tyr Tyr Cys Ala Ala Asp Trp Lys Tyr Trp Thr Cys Gly Ala Gln 40 Thr Gly Gly Tyr Phe Gly Gln Trp Gly Gln Gly Ala Gln Val Thr Val 100 105 110 Ser Ser 45 (2) INFORMATION FOR SEQ ID NO: 20: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 125 amino acids
(B) TYPE: amino acid 50 (C) STRANDEDNESS: single (D) TOPOLOGY: linear 55 (ii) MOLECULE TYPE: protein (vii) IMMEDIATE SOURCE: (B) CLONE: camel "heavy chain immunoglobulin" V-region (11) 60 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20: Gly Gly Ser Val Gln Ala Gly Gly Ser Leu Arg Leu Ser Cys Asn Val 65 Ser Gly Ser Pro Ser Ser Thr Tyr Cys Leu Gly Trp Phe Arg Gln Ala

	Pro	Gly	Arg 35	Glu	Arg	Glu	Gly	Val 40	Thr	Ala	Ile	Asn	Thr 45	Asp	Gly.	Ser
5	Ile	Ile 50	Tyr	Ala	Ala	Asp	Ser 55	Val	Lys	Gly	Arg	Phe 60	Thr	Ile	Ser	Gln
	Asp 65	Thr	Ala	Lys	Glu	Thr 70	Val	His	Leu	Gln	Met 75	Asn	Asn	Leu	Gln	Pro 80
10	Glu	qaA	Thr	Ala	Thr 85	Tyr	Tyr	Сув	Ala	Ala 90	Arg	Leu	Thr	Glu	Met 95	Gly
15	Ala	Cys	Asp	Ala 100	Arg	Trp	Ala	Thr	Leu 105	Ala	Thr	Arg	Thr	Phe 110	Ala	Tyr
	Asn	Tyr	Trp 115	Gly	Gln	Cly	Thr	Gln 120	Val	Thr	Val	Ser	Ser 125			
20	(2) INFO	RMAT	ION I	FOR S	SEQ 1	ID NO): 2	1:								
25	(i)	(A) (B) (C)	LEI TYI STI	NGTH: PE: & RANDI	: 114 amino EDNES	TERIS ami aci ss: s lines	ino a id sing:	acid	3							
	(ii)	MOLE	ECULI	TYP	PE: p	prote	ein									
30	(vii)					E: ≥1 "}	neavy	, cha	ain i	immus	oglo	bul:	in" (/-reg	jion	(13)
	(xi)	SEOU	JENCE	B DES	CRIE	PTION	1: SI	EO II	NO:	21:	:					
		_	_					-			•					
35	Gly 1				Glu 5	Ala						Leu	Ser	Сув	Thr 15	Ala
35 40	1	Gly	Ser	Val	5	Ala Ser	Gly	Gly	Ser	Leu 10	Arg				15	
	1 Ser	Gly	Ser	Val Val 20	5 Ser		Gly Met	Gly Ala	Ser Trp 25	Leu 10 Phe	Arg Arg	Gln	Val	Pro 30	15 Gly	Gln
	1 Ser Glu	Gly Gly Arg	Ser Tyr Glu 35	Val Val 20 Gly	5 Ser Val	Ser	Gly Met Phe	Gly Ala Val 40	Ser Trp 25 Gln	Leu 10 Phe Thr	Arg Arg Ala	Gln Asp	Val Asn 45	Pro 30 Ser	15 Gly Ala	Gln Leu
‡0 ‡5	Ser Glu Tyr Lys 65	Gly Gly Arg Gly 50 Asn	Ser Tyr Glu 35 Asp	Val Val 20 Gly Ser Leu	5 Ser Val Val Tyr	Ser Ala Lys Leu 70	Gly Met Phe Gly 55 Gln	Gly Ala Val 40 Arg	Ser Trp 25 Gln Phe Arg	Leu 10 Phe Thr Thr	Arg Arg Ala Ile Leu 75	Gln Asp Ser 60 Gln	Val Asn 45 His	Pro 30 Ser Asp	15 Gly Ala Asn Asp	Gln Leu Ala Thr
‡ 0	Ser Glu Tyr Lys 65	Gly Gly Arg Gly 50 Asn	Ser Tyr Glu 35 Asp	Val Val 20 Gly Ser Leu	5 Ser Val Val Tyr	Ser Ala Lys Leu	Gly Met Phe Gly 55 Gln	Gly Ala Val 40 Arg	Ser Trp 25 Gln Phe Arg	Leu 10 Phe Thr Thr	Arg Arg Ala Ile Leu 75	Gln Asp Ser 60 Gln	Val Asn 45 His	Pro 30 Ser Asp	15 Gly Ala Asn Asp	Gln Leu Ala Thr
‡0 ‡5	Ser Glu Tyr Lys 65	Gly Gly Arg Gly 50 Asn	Ser Tyr Glu 35 Asp Thr	Val 20 Gly Ser Leu	Ser Val Val Tyr Cys 85	Ser Ala Lys Leu 70	Gly Met Phe Gly 55 Gln Ala	Gly Ala Val 40 Arg Met	Ser Trp 25 Gln Phe Arg	Leu 10 Phe Thr Thr Asn Lys 90	Arg Ala Ile Leu 75 Asp	Gln Asp Ser 60 Gln Arg	Val Asn 45 His Pro	Pro 30 Ser Asp Asp	Gly Ala Asn Asp Trp 95	Gln Leu Ala Thr 80 Ala
‡0 ‡5	Ser Glu Tyr Lys 65	Gly Gly Arg Gly 50 Asn Val	Ser Tyr Glu 35 Asp Thr	val val 20 Gly Ser Leu Tyr	Ser Val Val Tyr Cys 85	Ser Ala Lys Leu 70 Ala	Gly Met Phe Gly 55 Gln Ala	Gly Ala Val 40 Arg Met	Trp 25 Gln Phe Arg Lys	Leu 10 Phe Thr Thr Asn Lys 90	Arg Ala Ile Leu 75 Asp	Gln Asp Ser 60 Gln Arg	Val Asn 45 His Pro	Pro 30 Ser Asp Asp	Gly Ala Asn Asp Trp 95	Gln Leu Ala Thr 80 Ala
‡0 ‡5	Ser Glu Tyr Lys 65 Gly Glu	Gly Gly Arg Gly 50 Asn Val Pro	Ser Tyr Glu 35 Asp Thr Tyr	Val 20 Gly Ser Leu Tyr Glu	Ser Val Val Tyr Cys 85 Trp	Ser Ala Lys Leu 70 Ala Asn	Gly Met Phe Gly 55 Gln Ala Asn	Gly Ala Val 40 Arg Met Gln Trp	Trp 25 Gln Phe Arg Lys	Leu 10 Phe Thr Thr Asn Lys 90	Arg Ala Ile Leu 75 Asp	Gln Asp Ser 60 Gln Arg	Val Asn 45 His Pro	Pro 30 Ser Asp Asp	Gly Ala Asn Asp Trp 95	Gln Leu Ala Thr 80 Ala
40 45 50	Ser Glu Tyr Lys 65 Gly Glu Ser (2) INFOR	Gly Arg Gly 50 Asn Val Pro Ser RMATI SEQU (A)	Ser Tyr Glu 35 Asp Thr Tyr Arg	Val 20 Gly Ser Leu Tyr Glu 100 FOR SECHAGTH:	Ser Val Val Tyr Cys 85 Trp GEQ I	Ser Ala Lys Leu 70 Ala Asn	Gly Met Phe Gly 55 Gln Ala Asn O: 22 STICS	Gly Ala Val 40 Arg Met Gln Trp	Trp 25 Gln Phe Arg Lys Gly 105	Leu 10 Phe Thr Thr Asn Lys 90	Arg Ala Ile Leu 75 Asp	Gln Asp Ser 60 Gln Arg	Val Asn 45 His Pro	Pro 30 Ser Asp Asp	Gly Ala Asn Asp Trp 95	Gln Leu Ala Thr 80 Ala

(ii) MOLECULE TYPE: protein

	(vii)	IMMI (B)	EDIAT	re so Dne:	OURCE	C: 21 "M	neavy	/ cha	in i	immu	noglo	bul:	in" '	V-re	gion	(16
5	(xi)	SEQ	JENCE	E DES	CRIE	OIT	1: SI	II Q	ON C	: 22	•					
.,	Gly 1	Gly	Ser	Ala	Gln 5	Ala	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Cys	Ala 15	Ala
10	His	Gly	Ile	Pro 20	Leu	Asn	Gly	Tyr	Tyr 25	Ile	Ala	Trp	Phe	Arg 30	Gln	Ala
	Pro	Gly	Lys 35	Gly	Arg	Glu	Gly	Val 40	Ala	Thr	Ile	Asn	Gly 45	Gly	Arg	Asp
15	Val	Thr 50	Tyr	Tyr	Ala	Asp	Ser 55	Vai	Thr	Gly	Arg	Phe 60	Thr	Ile	Ser	Arg
20	Asp 65	Ser	Pro	Lys	Asn	Thr 70	Val	Tyr	Leu	Gln	Met 75	Asn	Ser	Leu	Lys	Pro 80
	Glu	yab	Thr	Ala	Ile 85	Tyr	Phe	Cys	Ala	Ala 90	Gly	Ser	Arg	Phe	Ser 95	Ser
25	Pro	Val	Gly	Ser 100	Thr	Ser	Arg	Leu	Glu 105	Ser	Ser	Asp	Tyr	Asn 110	Tyr	Trp
	Gly	Gln	Gly 115	Ile	Gln	Val	Thr	Ala 120	Ser	Ser						
30	(2) INFO	RMAT:	ION I	FOR S	SEQ I	D NO): 23	3:				-				
35	(i)	(A (B (C	UENCE) LEM) TYI) STI) TOI	IGTH: PE: & RANDI	: 117 mino EDNES	am: ac: SS: 8	ino a id singl	cide	5							
40	(ii)	MOLI	ECULI	TYI	e: I	prote	ein									
-0	(vii)		EDIAT				neavy	, cha	ain i	immu	noglo	bul	in" '	V-reç	gion	(17
45	(xi)	SEQ	UENCI	DES	SCRIE	PTIO	v: SI	II QE	NO:	23:	:					
	Gly 1	Gly	Ser	Val	Gln 5	Pro	Gly	Gly	Ser	Leu 10	Thr	Leu	Ser	Cys	Thr 15	Val
50	Ser	Gly	Ala	Thr 20	Tyr	Ser	Asp	Tyr	Ser 25	Ile	Gly	Trp	Ile	Arg 30	Gln	Ala
	Pro	Gly	Lys 35	Asp	Arg	Glu	Val	Val 40	Ala	Ala	Ala	Asn	Thr 45	Gly	Ala	Thr
55	Ser	Lys 50	Phe	Tyr	Val	Asp	Phe 55	Val	Lys	Gly	Arg	Phe 60	Thr	Ile	Ser	Gln
60	Asp 65	Asn	Ala	Lys	Asn	Thr 70	Val	Tyr	Leu	Gln	Met 75	Ser	Phe	Leu	Lys	Pro 80
.,,	Glu	Asp	Thr	Ala	Ile 85	Tyr	Tyr	Cys	Ala	Ala 90	Ala	Asp	Pro	Ser	Ile 95	Tyr
65	Tyr	Ser	Ile	Leu 100	Xaa	Ile	Glu	Tyr	Lys 105	Tyr	Trp	Gly	Gln	Gly 110	Thr	Gln

Val Thr Val Ser Ser 115

5	(2) INFO	RMAT	ION I	FOR S	SEQ :	ID NO	D: 24	1:								
10	(i)	(B)	LEI TYI	NGTH: PE: & RANDE	12: mino EDNE:	3 am: 5 ac: 5S: !	ino a id sing!	acid	5							
	(ii)	MOLE	CULI	TY!	PE: 1	prote	∍in									
15	(Vii)						heavy	y cha	ain :	immu	noglo	bul:	in" '	V-re	gion	(18)
	(xi)	SEQU	JENCI	DES	CRI	PTIO	N: SI	EQ II	ON C	: 24:	:					
20	Gly 1	Gly	Ser	Val	Gln 5	Ala	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Cys	Thr 15	Gly
25	Ser	Gly	Phe	Pro 20	Tyr	Ser	Thr	Phe	Су s 25	Leu	Gly	Trp	Phe	Arg 30	Gln	Ala
	Pro	Gly	Lys 35	Glu	Arg	Glu	Gly	Val 40	Ala	Gly	Ile	Asn	Ser 45	Ala	Gly	Gly
30	Asn	Thr 50	Tyr	Tyr	Ala	Asp	Ala 55	Val	Lys	Gly	Arg	Phe 60	Thr	Ile	Ser	Gln
	Gly 65	Asn	Ala	Lys	Asn	Thr 70	Val	Phe	Leu	Gln	Met 75	Asp	Asn	Leu	Lys	Pro 80
3 5	Glu	yab	Thr	Ala	Ile 85	Tyr	Tyr	Сув	Ala	Ala 90	Asp	Ser	Pro	Сув	Tyr 95	Met
40	Pro	Thr	Met	Pro 100	Ala	Pro	Pro	Ile	Arg 105	Asp	Ser	Phe	Gly	Trp 110	Asp	Asp
40	Phe	Gly	Gln 115	Gly	Thr	Gln	Val	Thr 120	Val	Ser	Ser					
45	(2) INFO	RMAT:	ION I	FOR S	SEQ :	ID N	D: 2!	5:								
50	(i)	(B)) LEI	NGTH: PE: & RANDI	: 119 amine EDNE:	9 am: 5 ac: 55: :	ino a id sing:	acid	s							
	(ii)	MOLI	ECULI	E TYI	PE:	prot	≘in									
55	(Vii)	IMMI (B)	EDIA:	re so one:	OURC!	E: el "¦	heav	у съ	ain :	i m mu:	nogl	obul:	in" '	V-re	gion	(19)
	(xi)	SEQ	JENCI	E DES	SCRI	PTIO	N: 51	EQ II	ON O	: 25	:					
60	Gly 1	Gly	Ser	Val	Gln 5	Ala	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Cys	Ala 15	Ala
	Ser	Asp	Tyr	Thr 20	Ile	Thr	Asp	Tyr	Cys 25	Met	Ala	Trp	Phe	Arg 30	Gln	Ala
65	Pro	Gly	Lys 35	Glu	Arg	Glu	Leu	Val 40	Ala	Ala	Ile	Gln	Val 45	Val	Arg	Ser

	Asp	Thr 50	Arg	Leu	Thr	Asp	Tyr 55	Ala	Asp	Ser	Val	Lys 60	Gly	Arg	Phe.	Thr
5	Ile 65	Ser	Gln	Gly	Asn	Thr 70	Lys	Asn	Thr	Val	Asn 75	Leu	Gln	Met	Asn	Ser 80
	Leu	Thr	Pro	Glu	A sp 85	Thr	Ala	Ile	Tyr	Ser 90	Cys	Ala	Ala	Thr	Ser 95	Ser
10	Phe	Tyr	Trp	Tyr 100	Cys	Thr	Thr	Ala	Pro 105	Tyr	Asn	Val	Trp	Gly 110	Gln	Gly
15	Thr	Gln	Val 115	Thr	Val	Ser	Ser									
	(2) INFO	RMATI	ION 1	FOR S	SEO 1	D NO): 26	5 :								
20	(i)	(A) (B) (C)	LEI TYI STI	E CHANGTH: PE: 8 RANDE	: 117 Amino EDNES	ami aci	ino a id singl	cids	3							
25	(ii)	MOLI	ECULI	TYP	PE: p	rote	≥in									
	(vii)	IMMI (B)	EDIA:	re so one:	OURCE Came	C: el "1	neavy	y cha	ain i	Lmmur	noglo	bul:	in" \	/-rec	gion	(20)
30	(xi)	SEQ	JENCI	E DES	CRIE	PTIO	4: SI	II QE	NO:	26:	:					
	Gly 1	Gly	Ser	Val	Gln 5	Val	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Cys	Val 15	Ala
35	Ser	Thr	His	Thr 20	Asp	Ser	Ser	Thr	Cys 25	Ile	Gly	Trp	Phe	Arg 30	Gln	Ala
40	Pro	Gly	Lys 35	Glu	Arg	Glu	Gly	Val 40	Ala	Ser	Ile	Tyr	Phe 45	Gly	Asp	Gly
	Gly	Thr 50	Asn	Tyr	Arg	Asp	Ser 55	Val	Lys	Gly	Arg	Phe 60	Thr	Ile	Ser	Gln
45	Leu 65	Asn	Ala	Gln	Asn	Thr 70	Val	Tyr	Leu	Gln	Met 75	Asn	Ser	Leu	Lys	Pro 80
		•		Ala	85	-	-	-		90					95	
50	Gly	Сув	Asn	Leu 100	Arg	Thr	Thr	Phe	Thr 105	Arg	Trp	Gly	Gln	Gly 110	Thr	Gln
55	Val	Thr	Val 115	Ser	Ser											
	(2) INFO	RMAT:	ION	FOR S	SEQ :	D NO	o: 2	7:								
60	(i)	(A (B (C) LE	E CHANGTHE PE: A RANDI POLOG	: 125 amino EDNES	s am: c ac: ss: s	ino a id sing:	acid	5							
65	4223	MOT			DD											

	(Vii)						heav	y ch	ain	immu	nogl	obul	in"	V-re	gion	(21)
5	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: 51	EQ I	D NO	: 27	:					
•	Gly 1	Gly	Ser	Val	Gln 5	Val	Gly	Gly	Ser	Leu 10	Lys	Leu	Ser	Cys	Lys 15	Ile
10	Ser	Gly	Gly	Thr 20	Pro	Asp	Arg	Val	Pro 25	Lys	Ser	Leu	Ala	Trp 30	Phe	Arg
	Gln	Ala	Pro 35	Glu	Lys	Glu	Arg	Glu 40	Gly	Ile	Ala	Val	Leu 45	Ser	Thr	Lys
15	Asp	Gly 50	Lys	Thr	Phe	Tyr	Ala 55	Asp	Ser	Val	Lys	Gly Gly	Arg	Phe	Thr	Ile
20	Phe 65	Leu	Asp	Asn	Asp	Lys 70	Thr	Thr	Phe	Ser	Leu 75	Gln	Leu	Asp	Arg	Leu 80
	Asn	Pro	Glu	Asp	Thr 85	Ala	Asp	Tyr	Tyr	Cys 90	Ala	Ala	Asn	Gln	Leu 95	Ala
25 .				Tyr 100					105					Gly 110	Ala	Tyr
••	Ala	Ile	Trp 115	Gly	Gln	Gly	Thr	His 120	Val	Thr	Val	Ser	Ser 125			
30	(2) INFO	RMATI	ON I	FOR S	SEQ :	D NO): 2 8	3:								
35	(1)	(A) (B) (C)	LEI TYI STI	E CHI NGTH: PE: 8 RANDI POLOG	129 mino EDNES	ami aci	ino a id singl	cide	3							
40	(ii)	MOLE	ECULI	E TYI	PE: I	prote	≥in									
	(vii)						neavy	, cha	ain i	immur	noglo	buli	in" T	/-req	gion	(24)
45	(xi)	SEQU	JENCE	E DES	CRII	OIT	1: SI	EQ II	NO:	28:	:			•		
	Gly 1	Gly	Ser	Val	Gln 5	Ala	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Сув	Asn 15	Val
50	Ser	Gly	Ser	Pro 20	Ser	Ser	Thr	Tyr	Сув 25	Leu	Gly	Trp	Phe	Arg 30	Gln	Ala
55	Pro	Gly	Lys 35	Glu	Arg	Glu	Gly	Val 40	Thr	Ala	Ile	Asn	Thr 45	Asp	Gly	Ser
	Val	Ile 50	Tyr	Ala	Ala	Asp	Ser 55	Val	Lys	Gly	Arg	Phe 60	Thr	Ile	Ser	Gln.
60	Asp 65	Thr	Ala	Lys	Lys	Thr 70	Val	Tyr	Leu	Gln	Met 75	Asn	Asn	Leu	Gln	Pro 80
	Glu	Asp	Thr	Ala	Thr 85	Tyr	Tyr	Сув	Ala	Ala 90	Arg	Leu	Thr	Glu	Met 95	Gly
65	Ala	Cys	Asp	Ala 100	Arg	Trp	Ala	Thr	Leu 105	Ala	Thr	Arg	Thr	Phe 110	Ala	Tyr

Asn Tyr Trp Gly Arg Gly Thr Gln Val Thr Val Ser Ser 115 120 (2) INFORMATION FOR SEQ ID NO: 29: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 129 amino acids (B) TYPE: amino acid 10 (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein 15 (vii) IMMEDIATE SOURCE: (B) CLONE: camel "heavy chain immunoglobulin" V-region (25) (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29: 20 Gly Gly Ser Val Gln Thr Gly Gly Ser Leu Arg Leu Ser Cys Glu Ile Ser Gly Leu Thr Phe Asp Asp Ser Asp Val Gly Trp Tyr Arg Gln Ala 20 25 30 25 Gly Asp Glu Cys Lys Leu Val Ser Gly Ile Leu Ser Asp Gly Thr Pro Tyr Thr Lys Ser Gly Asp Tyr Ala Glu Ser Val Arg Gly Arg Val 50 60 30 Thr Ile Ser Arg Asp Asn Ala Lys Asn Met Ile Tyr Leu Gln Met Asn 65 - 70 75 80 35 Asp Leu Lys Pro Glu Asp Thr Ala Met Tyr Tyr Cys Ala Val Asp Gly
85 90 95 Trp Thr Arg Lys Glu Gly Gly Ile Gly Leu Pro Trp Ser Val Gln Cys
100 105 110 40 Glu Asp Gly Tyr Asn Tyr Trp Gly Gln Gly Thr Gln Val Thr Val Ser 115 120 125 Ser 45 (2) INFORMATION FOR SEQ ID NO: 30: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 111 amino acids (B) TYPE: amino acid 50 (C) STRANDEDNESS: single (D) TOPOLOGY: linear 55 (ii) MOLECULE TYPE: protein (vii) IMMEDIATE SOURCE: (B) CLONE: camel "heavy chain immunoglobulin" V-region (27) 60 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30: Gly Gly Ser Val Gln Ala Gly Gly Ser Leu Arg Leu Ser Cys Ala Ser 65 Ser Ser Lys Tyr Met Pro Cys Thr Tyr Asp Met Thr Trp Tyr Arg Gln 20 25 30

	Ala	Pro	Gly 35	Lys	Glu	Arg	Glu	Phe 40	Val	Ser	Ser	Ile	Asn 45	Ile	Asp	Gly
5	Lys	Thr 50	Thr	Tyr	Ala	Asp	Ser 55	Val	Lys	Gly	Arg	Phe 60	Thr	Ile	Ser	Gln
	Asp 65	Ser	Ala	Lys	Asn	Thr 70	Val	Tyr	Leu	Gln	Met 75	Asn	Ser	Leu	Lys	Pro 80
10	Glu	Asp	Thr	Ala	Met 85	Tyr	Tyr	Суѕ	Lys	Ile 90	Asp	Ser	Tyr	Pro	Cys 95	His
15	Leu	Leu	Asp	Val 100	Trp	Gly	Gln	Gly	Thr 105	Gln	Val	Thr	Val	Ser 110	Ser	
	(2) INFO	RMATI	ON E	FOR S	SEQ I	D NO): 3:	1:								
20	(i)	(A) (B) (C)	LEN TYI STI	E CHANGTH: PE: & RANDE	: 112 emino EDNES	2 ami o aci SS: s	ino a id singl	acids	5							
25	(ii)	MOLE	CULI	E TYP	E: p	prote	∍in									
·	(vii)						neavy	r cha	ain i	immur	noglo	bul:	in" (/-re	gion	(29)
30	(xi)	SEQU	JENCE	E DES	CRIE	OIT	N: SE	EQ II	NO:	31:	:					
	Gly 1	Gly	Ser -	Val	Gln 5	Ala	Gly	Gly	Ser	Leu 10	Arg	Leu	Ser	Сув	Val 15	Ala
35	Ser	Gly	Phe	Asn 20	Phe	Glu	Thr	Ser	Arg 25	Met	Ala	Trp	Tyr	Arg 30	Gln	Thr
40	Pro	Gly	Авп 35	Val	Сув	Glu	Leu	Val 40	Ser	Ser	Ile	Tyr	Ser 45	Asp	Gly	Lys
	Thr	Tyr 50	Tyr	Val	Asp	Arg	Met 55	Lys	Gly	Arg	Phe	Thr 60	Ile	Ser	Arg	Glu
45	Asn 65	Ala	Lys	Asn	Thr	Leu 70	Tyr	Leu	Gln	Leu	Ser 75	Gly	Leu	Lys	Pro	Glu 80
	Asp	Thr	Ala	Met	Tyr 85	Tyr	Cys	Ala	Pro	Val 90	Glu	Tyr	Pro	Ile	Ala 95	Asp
50	Met	Суѕ	Ser	Arg 100	Tyr	Gly	Asp	Pro	Gly 105	Thr	Gln	Val	Thr	Val 110	Ser	Ser
55	(2) INFO															
	(1)	(A)	LE	E CHA	: 416	bas	se pa									
60		(0)	STI	PE: 1 RANDI POLOC	EDNES	ss: s	sing	le								
	(ii)	MOLI	CULI	E TYI	PE: I	ANC	(gend	omic)							
<i>.</i> =	(vii)															
65		(B)	CL	ONE :			neavy FLAG					bul:	in" \	V-re	gion	follow

		(ix	•	A) N	AME/	KEY:		408										
5		(xi) SE	QUEN	CE DI	ESCR:	IPTI (ON:	SEQ :	ID N	D: 32	2:						
10						GAG Glu												48
10						TGT Cys												96
15						CCA Pro												144
20						ACC Thr												192
25						AAC Asn 70												240
30						GAC Asp												288
50						GGT Gly												336
35						GTC Val											٠.	384
40						GGT Gly		TAA:	TAGAI	ATT (3							416
45	(2)					SEQ CHAI				:								
50			(1	B) T	PE:	H: 13 amir DGY:	no a	cid	acio	is								
		-				YPE:	•											
						ESCRI												
55	Gln 1	Val	Lys	Leu	Leu 5	Glu	Ser	Gly	Gly	10	Ser	Val	GIn	Ala	15	GIA		
60	ser	Leu	Thr	Leu 20	Ser	Суз	Val	Tyr	Thr 25	Asn	Asp	Thr	Gly	Thr 30	Met	Gly		
	Trp	Phe	Arg 35	Gln	Ala	Pro	Gly	Lys 40	Glu	Сув	Glu	Arg	Val 45	Ala	His	Ile		
65	Thr	Pro 50		Gly	Met	Thr	Phe 55	Ile	Asp	Glu	Pro	Val 60	Lys	Gly	Arg	Phe		

	Thr 65	Ile	Ser	Arg	Asp	Asn 70	Ala	Gln	Lys	Thr	Leu 75	Ser	Leu	Arg	Met	Asn 80	
5	Ser	Leu	Arg	Pro	Glu 85	Asp	Thr	Ala	Val	Tyr 90	Tyr	Cys	Ala	Ala	Asp 95	Trp	
	Lys	Tyr	Trp	Thr 100	Cys	Gly	Ala	Gln	Thr 105	Gly	Gly	Tyr	Phe	Gly 110	Gln	Trp	
10	Gly	Gln	Gly 115	Ala	Gln	Val	Thr	Val 120	Ser	Ser	Leu	Ala	Ser 125	Tyr	Pro	Tyr	
15	Asp	Val 130	Pro	Asp	Tyr	Gly	Ser 135										
	(2)	INFO	ORMAT	rion	FOR	SEQ	ID 1	NO: 3	34:								
20		(i)	(2 (1 (0	QUENCA) LI B) TY C) ST O) TO	engti (Pe : [rani	nucl	3 ba leic ESS:	ase p acid	pairs 1	5							
25	-	(ii)	MOI	LECUI	E TY	PE:	DNA	(ger	nomic	=)							
30	((vii)		MEDIA B) CI		: can	mel '		-		immu			in"	V-re	egion	followed
,,		(ix)	(2	ATURI A) NI B) LO	ME/I			135									
35		(xi)	SEC	QUENC	CE DI	ESCRI	PTIC	ON: 8	SEQ I	ED NO	D: 34	4:					
10				CTG Leu													. 48
1 0				CTC Leu 20													96
1 5				TGG Trp													144
50				AAT Asn													192
5.5				GTG Val													240
. 0				TAT Tyr													288
50	ACG Thr	TAT Tyr	TAC Tyr	TGT Cys 100	GCG Ala	GCG Ala	GTC Val	CCA Pro	GCC Ala 105	CAC His	TTG Leu	GGA Gly	CCT Pro	GGC Gly 110	GCC Ala	ATT Ile	336
															GTC		

												CCG Pro 140					. 432
5	TAAT	LAGA!	ATT (C													443
	145																
10	(2)			TION									•				
15		•	` (<i>1</i> (1)	SEQUI A) LI B) TI D) TO	ENGTE (PE:	d: 14 amir	14 an	nino cid		_							
		(ii)	MOI	LECUI	LE TY	PE:	prot	ein									
20		(xi)	SE(QUENC	CE DI	ESCR	PTIC	ON: 8	SEQ :	ID NO): 3!	5:					
	Gln 1	Val	Lys	Leu	Leu 5	Glu	Ser	Gly	Gly	Gly 10	Ser	Val	Gln	Thr	Gly 15	Gly	
25	Ser	Leu	Arg	Leu 20	Ser	Сув	Ala	Val	Ser 25	Gly	Phe	Ser	Phe	Ser 30	Thr	Ser	
	Cys	Met	Ala 35	Trp	Phe	Arg	Gln	Ala 40	Ser	Gly	Lys	Gln	Arg 45	Glu	Gly	Val	• •
30	Ala	Ala 50	Ile	Asn	Ser	Gly	Gly 55	Gly	Arg	Thr	Tyr	Tyr 60	Asn	Thr	Tyr	Val	
35	Ala 65	Glu	Ser	Val-	Lys	Gly 70	Arg	Phe	Ala	Ile	Ser 75	Gln	Asp	Asn	Ala	Lys 80	
	Thr	Thr	Val	Tyr	Leu 85	Asp	Met	Asn	Asn	Leu 90	Thr	Pro	Glu	yab	Thr 95	Ala	
40	Thr	Tyr	Tyr	Cys 100	Ala	Ala	Val	Pro	Ala 105	His	Leu	Gly	Pro	Gly 110	Ala	Ile	
	Leu	Asp	Leu 115	Lys	Lys	Tyr	Lys	Tyr 120	Trp	Gly	Gln	Gly	Thr 125	Gln	Val	Thr	
45	Val	Ser 130	Ser	Leu	Ala	Ser	Tyr 135	Pro	Tyr	Asp	Val	Pro 140	Asp	Tyr	Gly	Ser	
50	(2)			TION QUEN		_											
55			(1	A) LI B) Ti C) Si D) To	(PE: [Rani	nuc: DEDNI	leic ESS:	acio	Ė	s							
		(ii	·	LECUI					omi	c)							
60	•	,	IMI	MEDIZ	ATE S	SOUR	CE:				immu	noglo	obul:	in" '	V-red	gion	followed
		(ix)) FE	ATURI	Ξ:	bу	the	FLAC	sec	quen	ce ()	pBŽ4)		•	-	
65		1 4 4 1	(I	A) NA B) LO	OCAT:	ION:	14		SEO '	ID NO	J. 34	۶.					

	CAG Gln l	GTG Val	AAA Lys	CTG Leu	CTC Leu 5	GAG Glu	TCT Ser	GGG Gly	GGA Gly	GGG Gly 10	TCG Ser	GTG Val	CAG Gln	GCT Ala	GGA Gly 15	GJA.	48
5	TCT Ser	CTG Leu	AGA Arg	CTC Leu 20	TCC Ser	TGT Cys	AAT Asn	GTC Val	TCT Ser 25	GGC Gly	TCT Ser	CCC Pro	AGT Ser	AGT Ser 30	ACT Thr	TAT Tyr	96
10	TGC Cys	CTG Leu	GGC Gly 35	TGG Trp	TTC Phe	CGC Arg	CAG Gln	GCT Ala 40	CCA Pro	GGG Gly	AAG Lys	GAG Glu	CGT Arg 45	GAG Glu	GGG Gly	GTC Val	144
15	ACA Thr	GCG Ala 50	ATT Ile	AAC Asn	ACT Thr	GAT Asp	GGC Gly 55	AGT Ser	GTC Val	ATA Ile	TAC Tyr	GCA Ala 60	GCC Ala	GAC Asp	TCC Ser	GTG Val	192
20	AAG Lys 65	GGC Gly	CGA Arg	TTC Phe	ACC Thr	ATC Ile 70	TCC Ser	CAA Gln	GAC Asp	ACC Thr	GCC Ala 75	AAG Lys	AAA Lys	ACG Thr	GTA Val	TAT Tyr 80	240
20	CTC Leu	CAG Gln	ATG Met	AAC Asn	AAC Asn 85	CTG Leu	CAA Gln	CCT Pro	GAG Glu	GAT Asp 90	ACG Thr	GCC Ala	ACC Thr	TAT Tyr	TAC Tyr 95	TGC Cys	288
25 .	GCG Ala	GCA Ala	AGA Arg	CTG Leu 100	ACG Thr	GAG Glu	ATG Met	GGG Gly	GCT Ala 105	TGT Cys	GAT Asp	GCG Ala	AGA Arg	TGG Trp 110	GCG Ala	ACC Thr	336
30	TTA Leu	GCG Ala	ACA Thr 115	AGG Arg	ACG Thr	TTT Phe	GCG Ala	TAT Tyr 120	AAC Asn	TAC Tyr	TGG Trp	GGC Gly	CGG Arg 125	Gly	ACC Thr	CAG Gln	384
35	GTC Val	ACC Thr 130	GTC Val	TCC Ser	TCA Ser	CTA Leu	GCT Ala 135	AGT Ser	TAC Tyr	CCG Pro	TAC Tyr	GAC Asp 140	GTT Val	ccs Pro	GAC Asp	TAC Tyr	432
40	GGT Gly 145		TAA	TAGA7	ATT C	2											449
	(2)	INFO	ORMAI	MOI	FOR	SEQ	ID N	10: 3	37:								
45		((<i>I</i>	1) LE 3) TY	NGTI PE:	i: 14 amir	RACTE 16 am no ac line	nino cid									
50		(11)	MOI	ECUI	E TY	PE:	prot	ein									
		(xi)	SEÇ	QUENC	E DE	SCRI	PTIC	on: S	EQ 1	D NO): 37	7 :					
55	Gln 1	Val	Lys	Leu	Leu 5	Glu	Ser	Gly	Gly	Gly 10	Ser	Val	Gln	Ala	Gly 15	Gly	
رر	Ser	Leu	Arg	Leu 20	Ser	Cys	Asn	Val	Ser 25	Gly	Ser	Pro	Ser	ser 30	Thr	Tyr	
60	Cys	Leu	Gly 35	Trp	Phe	Arg	Gln	Ala 40	Pro	Gly	Lys	Glu	Arg 45	Glu	Gly	Val	
	Thr	Ala 50	Ile	Asn	Thr	Asp	Gly 55	Ser	Val	Ile	Tyr	Ala 60	Ala	Asp	Ser	Val	
65	Lys 65	Gly	Arg	Phe	Thr	Ile 70	Ser	Gln	Asp	Thr	Ala 75	Lys	Lys	Thr	Val	Tyr 80	

	Leu	Gln	Met	Asn	Asn 85	Leu.	Gln	Pro	Glu	Asp 90	Thr	Ala	Thr	Tyr	Tyr 95	Сув		
5	Ala	Ala	Arg	Leu 100	Thr	Glu	Met	Gly	Ala 105	Сув	Asp	Ala	Arg	Trp 110	Ala	Thr		
	Leu	Ala	Thr 115	Arg	Thr	Phe	Ala	Tyr 120	Asn	Tyr	Trp	Gly	Arg 125	Gly	Ìhr	Gln		
10	Val	Thr 130	Val	Ser	Ser	Leu	Ala 135	Ser	Tyr	Pro	Tyr	Asp 140	Val	Pro	Asp	Tyr		
15	Gly 145	Ser																
	(2)	INF	ORMA'	TION	FOR	SEQ	ID I	NO:	38:									
20		(i	(1	A) Li B) T' C) S'	CE CI ENGTI YPE: TRANI OPOLO	H: 1 nuc DEDNI	19 ba leic ESS:	ase ; aci sin	pair: d	s								
25		(ii) MO	LECU	LE T	YPE:	DNA	(ge	nomi	C)								
•		(vii			ATE :			gure	6									
30		(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0: 3	8:						
	AAT	TTAG	CGG	CCGC	CCAG	GT G	AAAC	TGCT	C GA	GTAA	GTGA	CTA	AGGT	CAC	CGTC	TCCTC	A	60
35	GAA	CAAA	AAC	TCAT	CTCA	ga a	GAGG.	ATCT	g aa	TTAA	TGAG	AAT	TCAT	CAA :	ACGG'	TGATA		119
	(2)	INF	ORMA	TION	FOR	SEQ	ID	NO:	39:									
40		(i	· (A) L B) T C) S	CE C ENGT YPE: TRAN OPOL	H: 1 nuc DEDN	20 b leic ESS:	ase aci sin	pair d	S								
45		(ii) MO	LECU	LE T	YPE:	DNA	(ge	nomi	.c)								
		(vii			ATE LONE			gure	6			,						
50		(xi	.) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	ю: 3	9:						
	AGC	TTAT	CAC	CGTT	TGAT	GA A	TTCI	CATT	TA A'	TCAG	ATCC	TCI	TCTG	AGA	TGAG	TTTTI	rG	6
55	TTC	TGAG	GAG	ACGG	TGAC	CT T	AGTC	ACTT	'A CI	CGAG	CAGI	TTC	ACCT	GGG	cecc	CGCTA	VA.	120
	(2)	INF	ORMA	TION	FOR	SEÇ	ID	NO:	40:									
60		i)	· ((A) I (B) I (C) S	ICE C LENGT TYPE: TRAN	H: 7 ami DEDN	ami no a ESS:	no a cid sin	cids	3								
65		(ii	L) MC	DLECU	JLE 1	YPE:	pro	teir	1									

	(Vii) IMMEDIATE SOURCE: (B) CLONE: See figure 6	
5	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40:	
J	Ala Gln Val Lys Leu Glu 1 5	
10	(2) INFORMATION FOR SEQ ID NO: 41:	
15	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 16 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: protein	
20	(vii) IMMEDIATE SOURCE: (B) CLONE: See figure 6	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41:	
25	Val Thr Val Ser Ser Glu Gln Lys Leu Ile Ser Glu Glu Asp Leu Asn 1 5 10 15	
30	(2) INFORMATION FOR SEQ ID NO: 42:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 117 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
40	(vii) IMMEDIATE SOURCE: (B) CLONE: See figure 19	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 42:	
45	AATTTAGTCG CGACAGGTGA AACTGCTCGA GTAAGTGACT AAGGTCACCG TCTCCTCAGA	60
43	ACAAAAACTC ATCTCAGAAG AGGATCTGAA TTAATGAGAA TTCATCTTAA GGTGATA	117
50	(2) INFORMATION FOR SEQ ID NO: 43:	
55	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 117 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
60	<pre>(vii) IMMEDIATE SOURCE: (B) CLONE: See figure 19</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 43:	
65	AGCTTATCAC CTTAAGATGA ATTCTCATTA ATTCAGATCC TCTTCTGAGA TGAGTTTTTG	60
O.	TTCTGAGGAG ACGGTGACCT TAGTCACTTA CTCGAGCAGT TTCACCTGTC GCGACTA	117

```
(2) INFORMATION FOR SEQ ID NO: 44:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 6 amino acids
               (B) TYPE: amino acid
 5
               (C) STRANDEDNESS: single
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: protein
10
        (vii) IMMEDIATE SOURCE:
               (B) CLONE: See figure 19
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 44:
15
          Arg Gln Val Lys Leu Leu
20
     (2) INFORMATION FOR SEQ ID NO: 45:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 16 amino acids
               (B) TYPE: amino acid
(C) STRANDEDNESS: single
25
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: protein
30
        (vii) IMMEDIATE SOURCE:
               (B) CLONE: See figure 19
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 45:
35
          Val Thr Val Ser Ser Glu Gln Lys Leu Ile Ser Glu Glu Asp Leu Asn
                                                10
     (2) INFORMATION FOR SEQ ID NO: 46:
40
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 4 amino acids
               (B) TYPE: amino acid
               (C) STRANDEDNESS: single
45
               (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: protein
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 46:
50
          Gln Val Lys Leu
55
     (2) INFORMATION FOR SEQ ID NO: 47:
          (i) SEQUENCE CHARACTERISTICS:
               (A) LENGTH: 5 amino acids
               (B) TYPE: amino acid
60
               (C) STRANDEDNESS: single (D) TOPOLOGY: linear
         (ii) MOLECULE TYPE: protein
65
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 47:
```

	Val Thr Val Ser Ser 1 5	
5	(2) INFORMATION FOR SEQ ID NO: 48:	
10	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 48:	
	GTCACCGTCT CCTCATAATG A	2
20	(2) INFORMATION FOR SEQ ID NO: 49:	
25	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
30	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 49:	
	AGCTTCATTA TGAGGAGACG	20
3 5	(2) INFORMATION FOR SEQ ID NO: 50:	
40	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 34 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 50:	
	GTCACCGTCT CCTCATAATG ATCTTAAGGT GATA	34
50	(2) INFORMATION FOR SEQ ID NO: 51:	
55	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 33 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
60	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 51:	

65

AGCTTATCAC CTTAAGATCA TTATGAGGAG ACG

	(2) INFORMATION FOR SEQ ID NO: 52:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 12 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 52:	
	AATTGCGGCC GC	12
15	(2) INFORMATION FOR SEQ ID NO: 53:	
20	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 16 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
25	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 53:	
	CATGCAGTCT TCGGGC	16
30	(2) INFORMATION FOR SEQ ID NO: 54:	
35	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 16 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 54:	
	TTAAGCCCGA AGACTG	16
45	(2) INFORMATION FOR SEQ ID NO: 55:	
50	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 44 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
55	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 55:	
	TCACTGAATT CGGGATCATG AGGACTCTCC TTGTGAGCTC GCTT	44
60	(2) INFORMATION FOR SEQ ID NO: 56:	
65	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 48 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

	(11) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 56:	
5	ATGTCACAAA GCTTAAGCAC GAAGACAGTC GACCGTGCGG CCGGAGAC	48
	(2) INFORMATION FOR SEQ ID NO: 57:	
10	**	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 44 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
15	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 57:	
20	CGCGTCCATG CAGTCCTCAG GTGGATCATC CCAGGTGAAA CTGC	44
	(2) INFORMATION FOR SEQ ID NO: 58:	
25	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 44 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single	
30	(D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 58:	
35	TCGAGCAGTT TCACCTGGGA TGATCCACCT GAGGACTGCA TGGA	44
	(2) INFORMATION FOR SEQ ID NO: 59:	
40	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 15 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
45	(ii) MOLECULE TYPE: protein	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 59:	
50	Ser Met Gln Ser Ser Gly Gly Ser Ser Gln Val Lys Leu Leu Glu 1 5 10 15	
55	(2) INFORMATION FOR SEQ ID NO: 60:	
60	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 53 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: DNA (genomic)	
<i>(5</i>	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 60:	
65		

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	(2) INFORMATION FOR SEQ ID NO: 61:	
5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 53 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
10	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 61:	
	GGCCGCTGAG GAGACGGTGA CCTTAGTCAC TTACTCGAGC AGTTTCACCT GGC	53
15	(2) INFORMATION FOR SEQ ID NO: 62:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 6 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
25	(ii) MOLECULE TYPE: protein	
دع	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 62:	
30	Ser Ser Gly Gly Ser Ser	

30

CLAIMS

- A process for the production of an antibody or a fragment or functionalized fragment thereof using a transformed lower eukaryotic host containing an
 expressible DNA sequence encoding the antibody or (functionalized) fragment thereof, wherein the antibody or (functionalized) fragment thereof is derived from a heavy chain immunoglobulin of Camelidae and is devoid of light chains, and wherein the lower eukaryotic host is a mould or a yeast.
- 2. A process according to claim 1, in which the mould belongs to the genera Aspergillus or Trichoderma.
 - 3. A process according to claim 1, in which the yeast belongs to the genera Saccharomyces, Kluyveromcyes, Hansenula, or Pichia.
 - 4. A process according to claim 1, in which the heavy chain fragment at least contains the whole variable domain.
- A process according to claim 1, in which the antibody or (functionalized)
 fragment thereof derived from a heavy chain immunoglobulin of Camelidae comprises a complementary determining region (CDR) different from the CDR belonging to the natural antibody ex Camelidae grafted on the framework of the variable domain of the heavy chain immunoglobulin ex Camelidae.
- 6. A process according to claim 1, in which the immunoglobulin to be produced is a catalytic antibody raised in *Camelidae*.
 - 7. A process according to claim 1, in which the functionalized antibody or fragment thereof comprises a fusion protein of both a heavy chain immunoglobulin from *Camelidae* or a fragment thereof and another polypeptide.

8. A process according to claim 1, in which the DNA sequence encodes a modified heavy chain immunoglobulin or (functionalized) fragment thereof derived from *Camelidae* and being devoid of light chains, and is made by random or directed mutagenesis or both.

5

- 9. A process according to claim 8, in which the resulting immunoglobulin or (functionalized) fragment thereof is modified such that
 - it is better adapted for production by the host cell, or
 - it is optimized for secretion by the lower eukaryotic host into the fermentation medium, or
 - its binding properties (kon and kon) are optimized, or
 - its catalytic activity is improved, or
 - it has acquired a metal chelating activity, or
 - its physical stability is improved.

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10

- 10. A composition containing a product produced by a process as claimed in any one of claims 1-9.
- 11. New product obtainable by a process as claimed in any one of claims 1-9.
- 12. A composition containing a new product as claimed in claim 11.

.

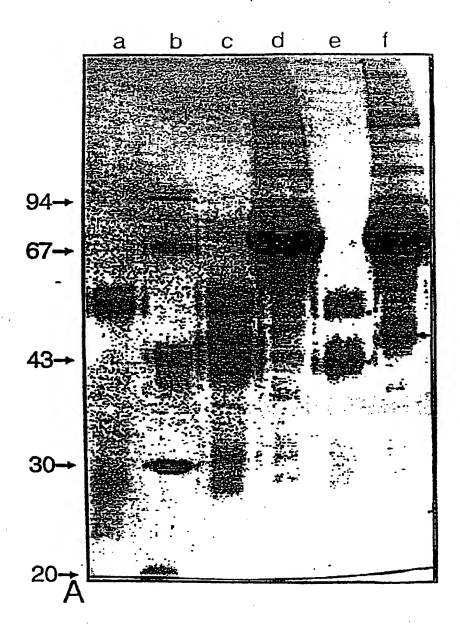
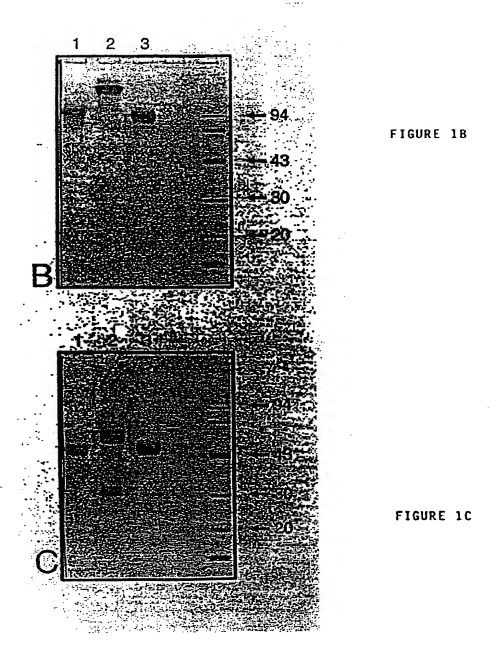
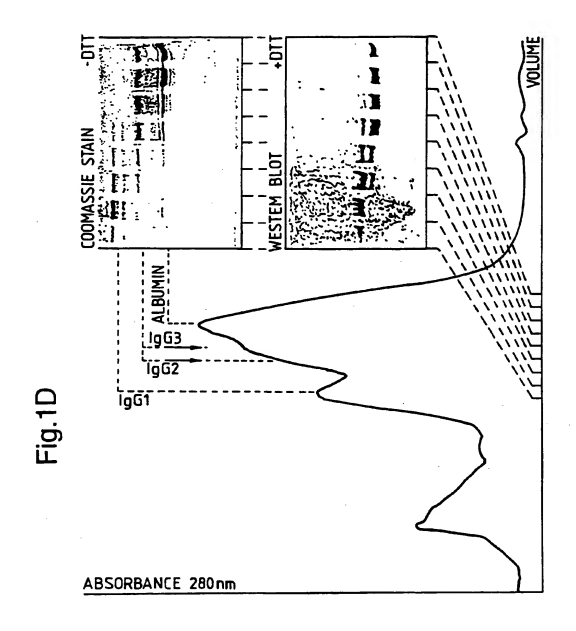


FIGURE 1A





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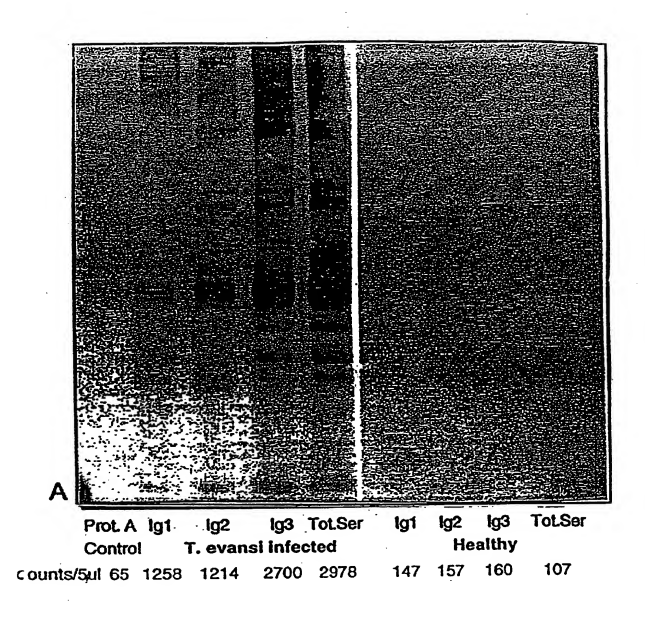


FIGURE 2A

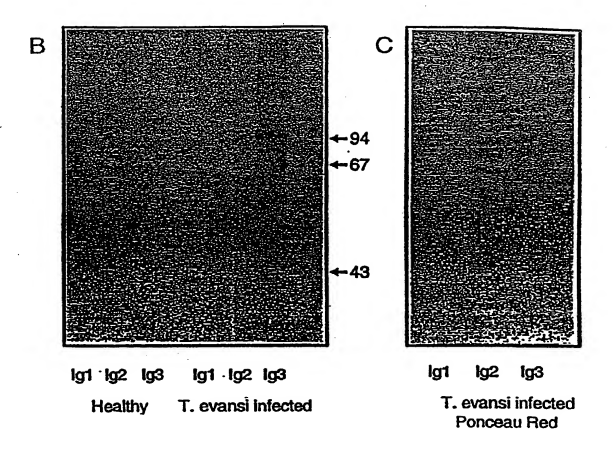


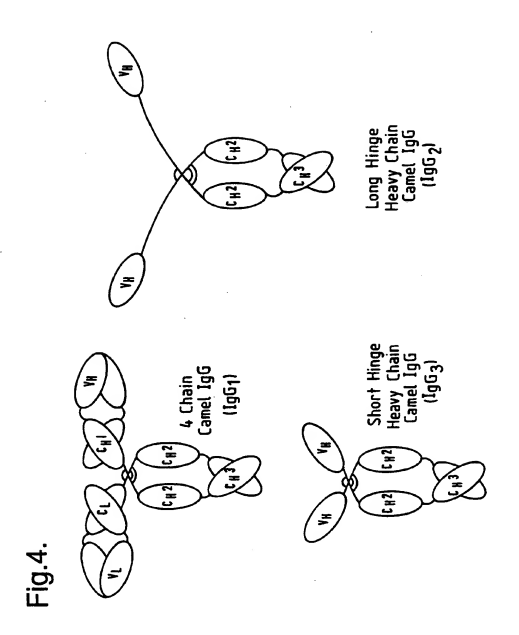
FIGURE 2B

FIGURE 2C

Fig.3.	20			40		
EVQLVESGGG	LVQPGGSLRL	SCAASG	CDR1	WVRQA	PGKGLEWVS	CDR2
GG	SVQGGGSLRL	SCAISG	CDR1	WFREG	PGKEREGIA	CDR2
GG	SVQAGGSLRL	SCASSS	CDR1	WYRQA	PGKEREFVS	CDR2

70	80				110	
RFTIS	RDNSKNTLYL	<i>QMISLRAEDTAVY</i>	YCAR	CDR3	WGQGTLVT	VSS
RFTIS	QDSTLKTMYL	LMNNLKPEDTGTY	YCAA	CDR3	WGQGTQVT	vss
RFTIS	QDSAKNTVYL	QMNSLKPEDTAMY	YCKI	CDR3	WGQGTQVT	vss

	came	ı v _H	hinge	C _H 2
camel	WGQGTQ	VT VSS	GTNEVCKCPKCP	APELPGG PSVFVFP
Camer	WGQGTQ	VT VSS	— EPKIPQPQPKPQPQP	
			OPOPKPOP	
			KPEPECTCPKCP	APELLGG PSVFIFP
•••••	human	C _H 1	hinge	C _H 2
human	gamma 3	KVDKRV	ELKTPLGDTTETCPRCP	•
			EPKCSDTPPPCPRCP	- • •
			EPKSCDTPPPCPRCP	APELLGG PSVFLFP
human	gamma 1	KVDKK-	AEPKSCDKTHTCPPCP	APELLGG PSVFLFP
human	gamma 2	KVKVTV	ERKCCVECPPCP	APPVAG - PSVFLFP
human	gamma 4	KVDKRV	ESKYGPPCPSCP	APEFLGG PSVFLFP



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Fig.5A.

,	CA	GGI	GAZ	VAC:	rcci	rcG/	AGT	CTG	GAG	GAG	CT	GG:	rgc	AGA(CTG	GAG	GAT	CTC	TGA	GACTO	60
•																		GAG.	ACT	CTGAC	
	Q	V	ĸ	L	L	Ε	S	G	G	G	s	V	Q	T	G	G	S	L	R	L	-
61				-+-							+			+-				+		AGGCT	120
																				rccga	•
	S	С	A	V	S	G	F	S	F	S	T	S	С	M	A	W	F	R	Q	A	-
121		AGG	AAA	GCA	GCG		GGG													ACTAC	180
•••		TCC	111	CGI	.cec															IGATG	
	s	G	K	Q	R	E	G	V	A	A	1	N	S	G	G	G	R	T	Y	Y	-
101																				CANG	240
101																				GITC	240
	N	T	¥	V	A	E	s	V	K	G	R	F	A	I	\$	Q	D	N	A	ĸ	-
241	AC	CAC	GGT																	CTCT	700
241	TG	GTG	CCA																	GACA	300
	T	T	V	Y	L	D	M	N	N	L	T	P	E	D	T	A	T	Y	· Y	С	-
		GGO	GGT	ccc											TI	GAA	AAA			.GTAC	
301		CCC	CCA	ccc			GAA				•				AAA	CII	111			CATC	360
	λ	A	V	P	A	R	L	G	P	G	λ	I	L	D	L	K	K	¥	ĸ	¥	-
	TG	GGG	CCA	GGG	GAC		Bst GGT			CIC	CTC	ACT	AGC	TAG	TTA	ccc	GTA	CGA	CGI	TCCG	
361				-+-			+				+			-+-							420
		G		G	T		v		v	s	s	L	A	s	Y	P	Y	D	v	P	_
421		CTA	ccc	TTC			Eco Gaa			•											
-41		GATY	SCC	AAG.			CII	AAG	44												

Fig.5B.

	CA	.GGT	GAA	ACI	Xh GCT	o I CGA	GTC	TGG	GGG	AGC	CTC	GGI	CA	GGG	TGO	GGG	GT	TCI	GAC	ACTC	
1		CCY			ACGAGCTCAGACCCCCTCCGAGCCACGTCCGACCCCCAGAGACTGTG		TGAG	60													
	Q	v	K	L	L	Ε	s	G	G	G	s	V	Q	Λ	G	G	S	L	T	L	-
											tyl Icol	•									
61	TC	TTG	TGT	ATA -+-	CAC	CVV	CGA	TAC	TGG	GAC	CAT	GGG	ATG	GT	TCC	:CC?	AGGC	TCC	AGG	GAAA	120
••	λG	ΛΛC	CACATATGTGGTTGCTATGACCCTGGTACCCTACCAAAGCGGTCCGAGG	TCC	CJ-L1.																
	s	C	v	Y	T	N	D	T	G	Т	M	G	W	F	R	Q	λ	P	G	K	-
																		TGA	ACC	CGTG	
121	CT	CAC	GCT	TTC	CÇA	ece	CGT	ATAATGCGGACTACCATACTGGAAGTAACTACTTGGGCAC	180												
	E	С	E	R	v	A	н	ı	T	P	D	G	M	T	F	I	D	E	P	v	-
81				-+-			+				+			-+-			+			GAAT	240
	TT	ccc	CCC	TAA	GTG	CTA	GAG	GGC	TCT	GTT	CCC	GGT	CII	TTG	CAA	CAG	AAA	CGC	TTA	CTTA	
	ĸ	G	R	F	T	I	s	R	D	N	A	Q	K	T	L	s	L	R	M	И	-
							CAC		CGT				TGC				GAA	ATA	CTG	GACT	300
41																	CII	TAT	GAC	CTGA	300
	s	L	R	P	E	D	T	A	v	Y	Y	С	A	A	D	W	ĸ	¥	W	Ŧ	-
	-	~ ~~	BCC	~~\	~ . ~	*			~~~	~~~		~~~		·				Bst		CGTC	
01				-+-			+				+			-+-			+			GCAG	360
																		v	T.	v	_
	С	G	A	Q	T	G	G	Y	F	G	Q	W	G	Q	G	λ	Q	-	_	•	_
61				-+-			+				+		CTA	-+-			+		TTC	416	
	λG	GAG	TGA	TCG	ATC	AAT	GGG	CAT	CCT	GCA	AGG	CCI	GAT	GCC	AA G	AAT	TAT	CIT	ХAG		
	5	S	T.	A	S	v	P	v	ח	v	P	D	Y	G	S	•	•				

Fig.5C.

	XhoI CAGGTGAAACTGCTCGAGTCTGGGGGGGGGGTCTCTGAGACTC GTCCACTTTGACGAGCTCAGACCCCCTCCCAGCCACGTCCGAGACCTCCCAGAGACTCTGAG	ACTC	60																		
1	GT	CCA	CTT	TGA	CGA	GCT	CAG	ACC	CCC	TCC	CAG	CCA	CGT	ccc	ACC	TCC	CAG	AGA	CTC	TGAG	80
	Q	v	ĸ	Ĺ	L	E	s	G	G	G	s	v	Q	A	G	G	5	L	R	L	-
61							+				+			-+-			+			GGCT	120
				ACA: V				•	S								F		0	CCGA	_
		C	N	V	:>	(,	5	r	3	3	1	•		L	Ů	••	•	••	•		
121	CC	AGG	GAA	GGA	ece.	TGA	GGG +	GGT	CAC	AGC	GAT	TAA	CAC	TGA	TGG	CAG	TGT +	CAT	ATA	CGCA	180
	GG'	TCC	CTT	CCT	CGC	ACT	ccc	CCA	.GTG	TCG	CTA	ATT	GTG	ACT	ACC	GIC	ACA	GTA	TAT	GCGT	
	P	G	ĸ	E	R	E	G	V	T	λ	I	И	T	D	G	S	V	I	Y	A	-
181	GC	CGA	CIC	CGT	Gλλ	GGG	cce	ATT	CVC	CAT	CTC	CCA	AGA	CAC	CGC	CVV	+ Gእአ	λλC	GGT.	ATAT	240
101	CG	CT	GAG	GCA	CIT	CCC	GGC	TAA	GTG	GTA	GAG	GGT	TCT	GTG	GCG	GTT	CIT	TTG	CCY.	ATAT	
	A	D	S	V	К.	G	R	F	T	I	S	Q	D	T	A	K	K	T	v	Y	-
241		CCA	GAT																AAG.	ACTG	30 0
	GΛ	GGT	CTA																	I GYC	
	L	Q	M	N	И	L	Q	P	E	D	T	λ	T	Y	Y	С	γ	λ	R	L	-
301	AC	GGA	GAT	GGG	GGC	TTG	ΤGλ	TGC	GAG	ATG	GGC	GAC	CTT	AGC	GAC	AAG	GAC	GTT	TGC	GTAT	360
301	TG	CCI	CTA	ccc	CCG	AAC	ACT	'ACG	CTC	TAC	coc	CIG	GAA	TOG	CIG	TTC	CIG	CAA	ACG	CATA	
	T	E	M	G	A	C	D	λ	R	W	λ	T	L	A	T	R	T	F	A	¥	-
	AA	CTA	CTG	GGG	CCG	GGG	GAC	CCA		EII		CIC	CTC	:XCI	AGC	TAG	TTA	.ccc	GTA	CGAC	
361				-+-			+				+			-+-			+			ecre	420
	N	Y	W	G	R	G	T	Q	v	T	v	s	s	L	λ	s	Y	P	¥	D	-
427		TCC	GGA	CTA	CGG	TTC			ECC	TTC		۵.									
421		AGG	CCT	GAT	GCC	AAG						. 3									
		_	_	3.5	_	_	_	_													

9 (ECORI) EagI AATTTAGCGCCCCCAGGTGAAACTGCTCGAGTAAGTCACTAAAGGTCACCGTCTCCTCA AATCGCCGGCGGGTCCACTTTGACGAGCTCATTCACTGATTCCAGTGGCAGAGGAGT

CTTGTTTTTGAGTAGAGTCTTCTCCTAGACTTAATTACTCTTAAGTAGTTTGCCACTATT

E Q K L I S E E D L N * * GAACAAAAACTCATCTCAGAAGAGGATCTGAATTAATGAGAATTCATCAAACGGTGATA ECORI

61

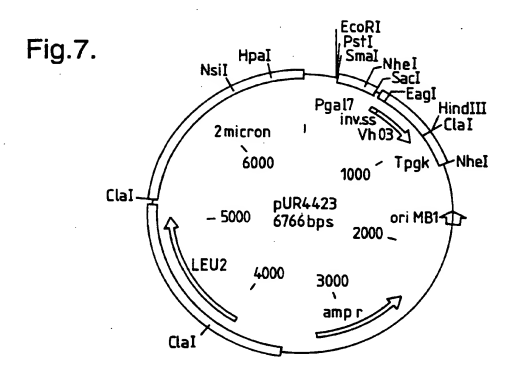
HindIII

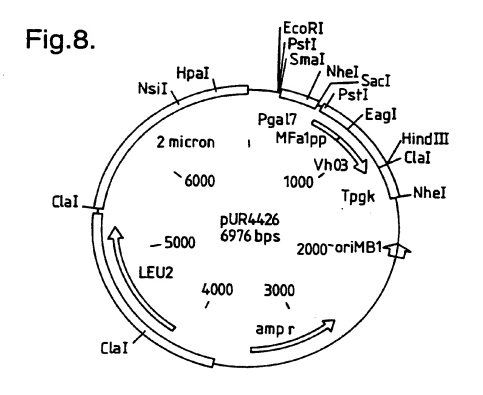
--- 123 CGA 121

9 ECORI) NEUI XHOI BSEEII
AATTTAGICGCGACAGGIGAAACIGCTCGAGAAGTGACTAAGGICACGGTCTCCCTCAGA ATCAGCGCTGTCCACTTGAGGAGCTCATTCACTGATTCCAGTGGAGGAGTCT R Q V K L L V S S E (EcoRI) NruI

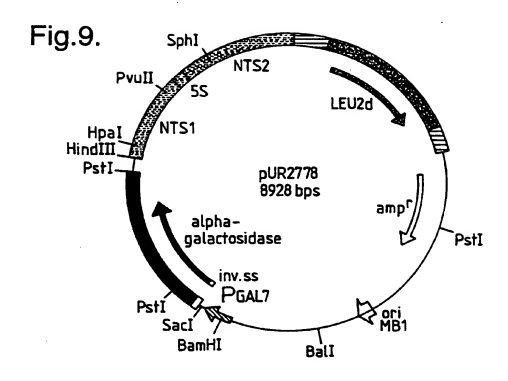
HindIII TCTTTTTCAGTAGACTCTTCTCCTAGACTTAATTACTCTTAAGTAGAATTCCACTATTCG
Q K L I S E E D L N * * Ecori actiti ni 61

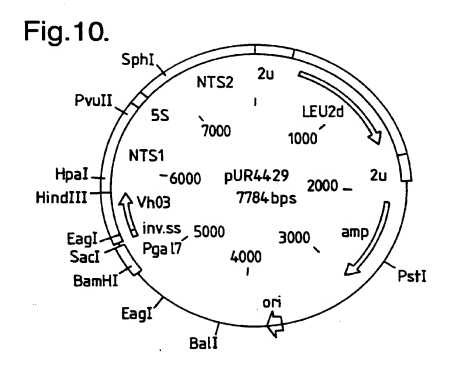
- 121 A 121





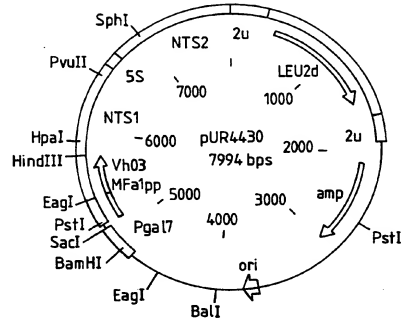
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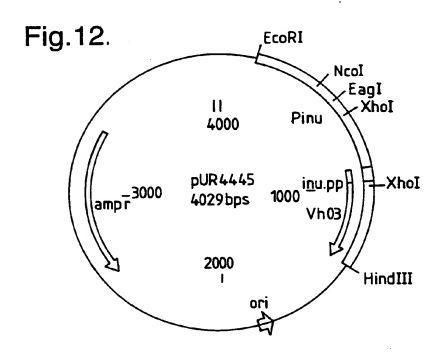


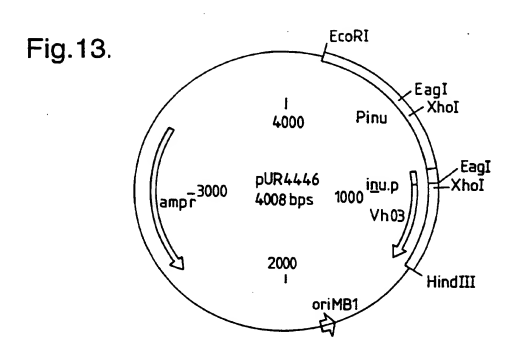


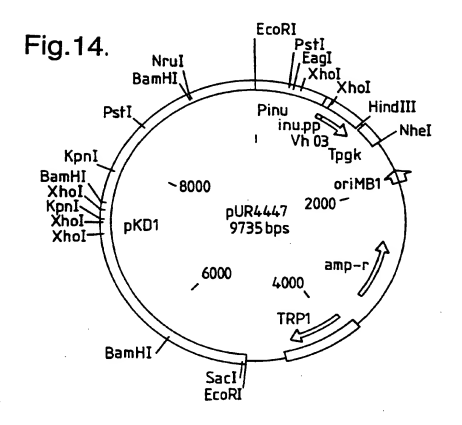
SUBSTITUTE SHEET (RULE 26)

Fig.11.



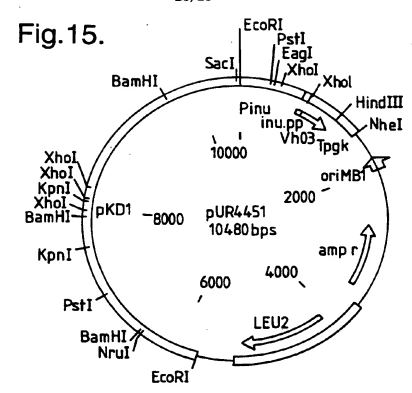


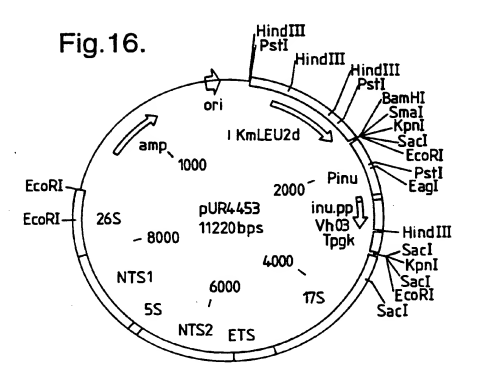




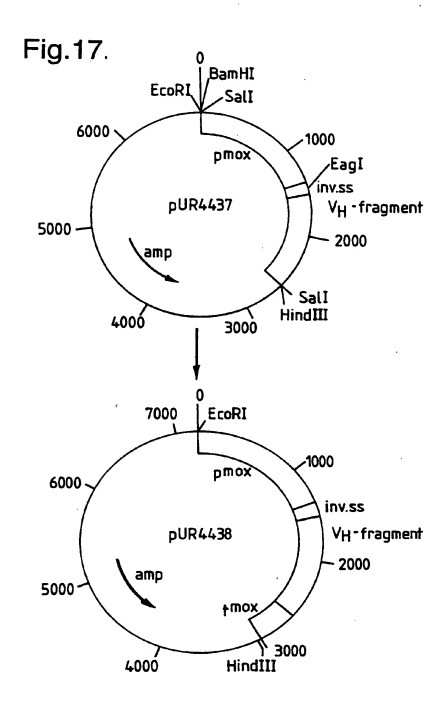
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Fig. 18.

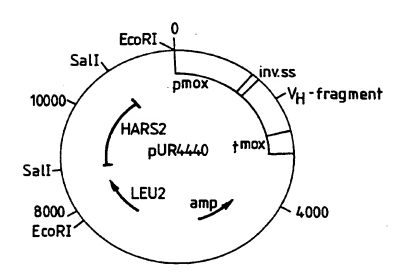
9000 | EcoRI | 1000 |

inv.ss | pmox | 2000 | EcoRI |

7000 | LEU2 | 3000 |

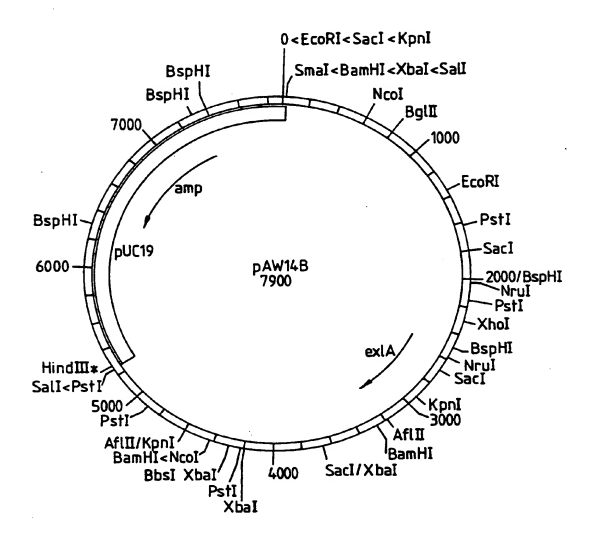
ÈcoRI

6000



4000

Fig.20.



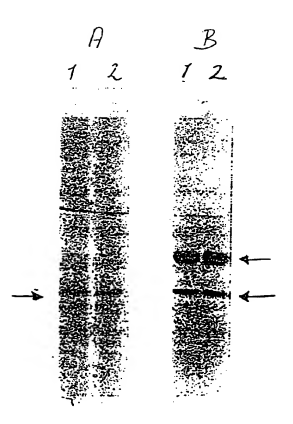


Figure 21

INTERNATIONAL SEARCH REPORT

Inter nal Application No PCT/EP 94/01442

A. CLASSI IPC 5	FICATION OF SUBJECT MATTER C12N15/13 C07K15/28 A61K39/	/395	
According to	o International Patent Classification (IPC) or to both national class	sification and IPC	
	SEARCHED		
Minimum de IPC 5	ocumentation searched (classification system followed by classific C12N C07K A61K	ation symbols)	
Documentati	ion searched other than minimum documentation to the extent tha	t such documents are included in the fields	tearched
Electronic d	ata base consulted during the international search (name of data b	ase and, where practical, search terms used)	
C. DOCUM	IENTS CONSIDERED TO BE RELEVANT		
Category '	Citation of document, with indication, where appropriate, of the	relevant passages	Relevant to claim No.
A	EP,A,O 256 421 (PHILLIPS PETROLE COMPANY) 24 February 1988 cited in the application see the whole document	EUM	1,3
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X Furt	her documents are listed in the continuation of box C.	Y Patent family members are listed	in annex.
* Special car	tegories of cited documents:	erre land de manage ambléhad after the ins	and all Glips date
'A' docum consid 'E' earlier filing ('L' docum which citation	ent defining the general state of the art which is not ered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another n or other special reason (as specified) ent referring to an oral disclosure, use, exhibition or	"T" later document published after the init or priority date and not in conflict we cited to understand the principle or to invention. "X" document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the document of particular relevance; the cannot be considered to involve an indocument is combined with one or in ments, such combination being obvi	ith the application but heory underlying the claimed invention t be considered to occurrent is taken alons claimed invention when the hore other such docu-
'P' docum	ent published prior to the international filing date but han the priority date claimed	in the art. "&" document member of the same paters	-
Date of the	actual completion of the international search	Date of mailing of the international s	/ ·
1	9 August 1994	2 6 -08- 1994	1
Name and s	mailing address of the ISA European Patent Office, P.B. 5818 Patentlasn 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Far (+31-70) 340-3016	Authorized officer Nooij, F	

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INTERNATIONAL SEARCH REPORT

Inte mai Application No
PCT/EP 94/01442

		PC1/EP 94/U1442
(Continue	tion) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
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P,X	FEBS LETTERS vol. 339, no. 3 , 21 February 1994 , AMSTERDAM, THE NETHERLANDS pages 285 - 290 J. DAVIES ET AL. ''Camelising' human antibody fragments: NMR studies on VH domains.' see the whole document	1,5, 10-12
P,X	WO,A,94 04678 (C. CASTERMAN ET AL.) 3 March 1994 see the whole document	1,3,4,6, 10-12
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